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INTERLEUKIN-17 RELATED MAMMALIAN CYTOKINE (IL-171). POLYNUCLEOTIDES ENCODING THEM. USES

This application is a PCT filing claiming priority to U.S. Patent Application USSN 09/229,402, filed January 11, 5 1999.

### FIELD OF THE INVENTION

The present invention relates to compositions related to proteins which function in controlling physiology, development, and differentiation of mammalian cells, e.g., 10 cells of a mammalian immune system. In particular, it provides nucleic acids, proteins, antibodies, and mimetics which regulate cellular physiology, development, differentiation, or function of various cell types, including 15 hematopoietic cells.

## BACKGROUND OF THE INVENTION

The immune system of vertebrates consists of a number of organs and several different cell types. Two major cell types include the myeloid and lymphoid lineages. Among the lymphoid 20 cell lineage are B cells, which were originally characterized as differentiating in fetal liver or adult bone marrow, and T cells, which were originally characterized as differentiating in the thymus. See, e.g., Paul (ed. 1998) Fundamental Immunology (4th ed.) Raven Press, New York. 25

In many aspects of the development of an immune response or cellular differentiation, soluble proteins known as cytokines play a critical role in regulating cellular These cytokines apparently mediate cellular activities in many ways. They have been shown, in many cases, to modulate proliferation, growth, and differentiation of hematopoietic stem cells into the vast number of progenitors composing the lineages responsible for an immune response.

However, the cellular molecules which are expressed by different developmental stages of cells in these maturation pathways are still incompletely identified. Moreover, the

roles and mechanisms of action of signaling molecules which induce, sustain, or modulate the various physiological, developmental, or proliferative states of these cells is poorly understood. Clearly, the immune system and its response to various stresses had relevance to medicine, e.g., infectious diseases, cancer related responses and treatment, allergic and transplantation rejection responses. See, e.g., Thorn, et al. <a href="Harrison's Principles of Internal Medicine">Harrison's Principles of Internal Medicine</a> McGraw/Hill, New York.

Medical science relies, in large degree, to appropriate recruitment or suppression of the immune system in effecting cures for insufficient or improper physiological responses to environmental factors. However, the lack of understanding of how the immune system is regulated or differentiates has blocked the ability to advantageously modulate the normal defensive mechanisms to biological challenges. Medical conditions characterized by abnormal or inappropriate regulation of the development or physiology of relevant cells thus remain unmanageable. The discovery and characterization of specific cytokines will contribute to the development of therapies for a broad range of degenerative or other conditions which affect the immune system, hematopoietic cells, as well as other cell types. The present invention provides solutions to some of these and many other problems.

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#### SUMMARY OF THE INVENTION

The present invention is based, in part, upon the discovery of cDNA clones encoding various cytokine-like proteins which exhibit significant sequence similarity to the cytokine designated CTLA-8.

The invention embraces isolated genes encoding the proteins of the invention, variants of the encoded proteins, e.g., mutations (muteins) of the natural sequences, species and allelic variants, fusion proteins, chemical mimetics, antibodies, and other structural or functional analogs. Various uses of these different nucleic acid or protein compositions are also provided.

In certain nucleic acid embodiments, the invention provides an isolated or recombinant polynucleotide comprising sequence from: a) a mammalian IL-171, which: encodes at least 8 contiguous amino acids of SEQ ID NO: 3 or 5; encodes at least two distinct segments of at least 5 contiguous amino acids of SEQ ID NO: 3 or 5; or comprises one or more segments at least 21 contiguous nucleotides of SEQ ID NO: 1 or 4; or b) a mammalian IL-175, which: encodes at least 8 contiguous amino acids of SEQ ID NO: 8; encodes at least two distinct segments 10 of at least 5 contiguous amino acids of SEO ID NO: 8; or comprises one or more segments at least 21 contiguous nucleotides of SEO ID NO: 6. Other embodiments include such a polynucleotide in an expression vector, comprising sequence: a) (IL-171) which: encodes at least 12 contiguous amino acids 15 of SEO ID NO: 3 or 5; encodes at least two distinct segments of at least 7 and 10 contiguous amino acids of SEQ ID NO: 3 or 5; or comprises at least 27 contiguous nucleotides of SEQ ID NO: 1 or 4; or b) (IL-175) which: encodes at least 12 contiguous amino acids of SEO ID NO: 8; encodes at least two 20 distinct segments of at least 7 and 10 contiguous amino acids of SEQ ID NO: 8; or comprises at least 27 contiguous nucleotides of SEO ID NO: 6. Certain embodiments will include those polynucleotides: a) (IL-171) which: encode at least 16 contiguous amino acids of SEQ ID NO: 3 or 5; encode at least two distinct segments of at least 10 and 14 contiguous amino 25 acid residues of SEQ ID NO: 3 or 5; comprise at least 33 contiguous nucleotides of SEQ ID NO: 1 or 4; or comprise SEQ ID NO: 1 or 4; or b) (IL-175) which: encode at least 16 contiguous amino acid residues of SEQ ID NO: 8; encode at 30 least two distinct segments of at least 10 and 13 contiguous amino acid residues of SEQ ID NO: 8; or comprise at least 33 contiguous nucleotides of SEQ ID NO: 6.

Various methods are provided, e.g., making: a) a polypeptide comprising expressing the described expression vector, thereby producing the polypeptide; b) a duplex nucleic acid comprising contacting a described polynucleotide with a complementary nucleic acid, thereby resulting in production of

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the duplex nucleic acid; or c) a described polynucleotide comprising amplifying using a PCR method.

Alternatively, the invention provides an isolated or recombinant polynucleotide which hybridizes under stringent wash conditions of at least 55° C and less than 400 mM salt to: a) the described (IL-171) polynucleotide which consists of SEQ ID NO: 1 or 4; or b) the described (IL-175) polynucleotide which consists of SEQ ID NO: 6 or 7. Other embodiments include such described polynucleotide: a) wherein the wash conditions are at least 65° C and less than 300 mM salt; or b) which comprises at least 50 contiguous nucleotides of the coding portion of: SEQ ID NO: 1 or 4 (IL-171); or SEQ ID NO: 6 or 7 (IL-175).

Certain kits are provided, e.g., comprising a described polynucleotide, and: a) instructions for the use of the polynucleotide for detection; b) instructions for the disposal of the polynucleotide or other reagents of the kit; or c) both a and b.

Various cells are provided also, e.g., a cell containing the described expression vector, wherein the cell is: a prokaryotic cell; a eukaryotic cell; a bacterial cell; a yeast cell; an insect cell; a mammalian cell; a mouse cell; a primate cell; or a human cell.

Polypeptide embodiments include, e.g., an isolated or recombinant antigenic polypeptide: a) (IL-171) comprising at least: i) one segment of 8 identical contiguous amino acids from SEQ ID NO: 3 or 5; or ii) two distinct segments of at least 5 contiguous amino acids from SEQ ID NO: 3 or 5; or b) (IL-175) comprising at least: i) one segment of 8 identical contiguous amino acids from SEQ ID NO: 8; or ii) two distinct segments of at least 5 contiguous amino acids from SEQ ID NO: 8. Additional embodiments include such a described polypeptide, wherein: a) the segment of 8 identical contiguous amino acids is at least 14 contiguous amino acids; or b) one of the segments of at least 5 contiguous amino acids comprises at least 7 contiguous amino acids. Other embodiments include a described polypeptide, wherein: A) (IL-171) the polypeptide:

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a) comprises SEQ ID NO: 3 or 5; b) binds with selectivity to a polyclonal antibody generated against an immunogen of SEQ ID NO: 3 or 5; c) comprises a plurality of distinct polypeptide segments of 10 contiguous amino acids of SEQ ID NO: 3 or 5; d) is a natural allelic variant of SEQ ID NO: 3 or 5; e) has a length at least 30 amino acids; or f) exhibits at least two non-overlapping epitopes which are selective for SEQ ID NO: 3 or 5; or B) (IL-175) the polypeptide: a) comprises SEQ ID NO: 8; b) binds with selectivity to a polyclonal antibody generated against an immunogen of SEQ ID NO: 8; c) comprises a 10 plurality of distinct polypeptide segments of 10 contiguous amino acids of SEQ ID NO: 8; d) has a length at least 30 amino acids; or e) exhibits at least two non-overlapping epitopes which are selective for SEQ ID NO: 8. Various other embodiments include such a described polypeptide, which: a) is 15 in a sterile composition; b) is not glycosylated; c) is denatured; d) is a synthetic polypeptide; e) is attached to a solid substrate; f) is a fusion protein with a detection or purification tag; g) is a 5-fold or less substitution from a natural sequence; or h) is a deletion or insertion variant 20 from a natural sequence.

Methods of using described polypeptides are also provided, e.g.,: a) to label the polypeptide, comprising labeling the polypeptide with a radioactive label; b) to separate the polypeptide from another polypeptide in a mixture, comprising running the mixture on a chromatography matrix, thereby separating the polypeptides; c) to identify a compound that binds selectively to the polypeptide, comprising incubating the compound with the polypeptide under appropriate conditions; thereby causing the compound to bind to the polypeptide; or d) to conjugate the polypeptide to a matrix, comprising derivatizing the polypeptide with a reactive reagent, and conjugating the polypeptide to the matrix.

Antibodies are also provided, including a binding

35 compound comprising an antigen binding portion from an
antibody which binds with selectivity to such a described
polypeptide, wherein the polypeptide: a) (IL-171) comprises

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SEQ ID NO 3 or 5; or b) (IL-175) comprises SEQ ID NO 8. Certain embodiments embrace such a binding compound, wherein the antibody is a polyclonal antibody which is raised against:

a) (IL-171) of SEQ ID NO: 3 or 5; or b) (IL-175) SEQ ID NO: 8. Other embodiments include such a described binding compound, wherein the: a) antibody: i) is immunoselected; ii) binds to a denatured protein; or iii) exhibits a Kd to the polypeptide of at least 30 mM; or b) the binding compound: i) is attached to a solid substrate, including a bead or plastic membrane; ii) is in a sterile composition; or iii) is detectably labeled, including a radioactive or fluorescent label.

Methods are provided, e.g., producing an antigen:antibody complex, comprising contacting a polypeptide comprising sequence from SEQ ID NO: 3, 5, or 8 with a described binding compound under conditions which allow the complex to form. Preferably, the binding compound is an antibody, and the polypeptide is in a biological sample.

Kits are provided, e.g., comprising a described binding compound and: a) a polypeptide of SEQ ID NO: 3, 5, or 8; b) instructions for the use of the binding compound for detection; or c) instructions for the disposal of the binding compound or other reagents of the kit.

And a method if provided of evaluating the selectivity of binding of an antibody to a protein of SEQ ID NO: 3, 5, or 8, comprising contacting a described antibody to the protein and to another cytokine; and comparing binding of the antibody to the protein and the cytokine.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

#### I. General

The present invention provides DNA sequence encoding various mammalian proteins which exhibit structural features characteristic of cytokines, particularly related to the cytokine designated CTLA-8 (also referred to as IL-17). Rat, mouse, human forms and a viral homolog of the CTLA-8 have been described and their sequences available from GenBank. See

Rouvier, et al. (1993) J. Immunol. 150:5445-5456; Yao, et al. (1995) <u>Immunity</u> 3:811-821; Yao, et al. (1995) <u>J. Immunol.</u> 155:5483-5486; and Kennedy, et al. (1996) J. Interferon and Cytokine Res. 16:611-617. The CTLA-8 has activities implicated in arthritis, kidney graft rejection, 5 tumorigenicity, virus-host interactions, and innate immunity; and appears to exhibit certain regulatory functions similar to IL-6. See PubMed (search for IL-17); Chabaud, et al. (1998) J. Immunol. 63:139-148; Amin, et al. (1998) Curr. Opin. Rheumatol. 10:263-268; Van Kooten, et al. (1998) J. Am. Soc. 10 Nephrol. 9:1526-1534; Fossiez, et al. (1998) Int. Rev. <u>Immunol.</u> 16:541-551; Knappe, et al. (1998) <u>J. Virol.</u> 72:5797-5801; Seow (1998) Vet. Immuno. Immunopathol. 63:139-48; and Teunissen, et al. (1998) <u>J. Invest. Dermatol.</u> 111:645-649. A report on the signaling through the NFKB transcription factor 15 implicates a signal pathway which is used in innate immunity. Shalom-Barak, et al. (1998) <u>J. Biol. Chem.</u> 273:27467-27473.

The newly presented cDNA sequences exhibit various features which are characteristic of mRNAs encoding cytokines, growth factors, and oncogenes. Because the IL-17 is the first 20 member of this newly recognized family of cytokines related to TGF- $\beta$ , Applicants have designated the family IL-170, with the new members IL-171 and IL-175; and IL-172, IL-173, IL-174, IL-176, and IL-177. The fold for this family is predicted to be that of the TGF- $\beta$  family of cytokines. The TGF- $\beta$  family of 25 cytokines, and the IL-170 family share the common feature of a cystine knot motif, characterized by a particular spacing of cysteine residues. See, e.g., Sun and Davies (1995) Ann. Rev. Biophys. Biomolec. Struct. 24:269-291; McDonald, et al. (1993) Cell 73:421-424; and Isaacs (1995) Curr. Op. Struct. Biol. 30 In particular, the structures suggest a number of 5:391-395. conserved cysteines, which correspond to, and are numbered, in human IL-172 (SEQ ID NO: 10), cysteines at 101, 103, 143, 156, and 158. The first cysteine corresponds to the postion in Table 7 of human IL-175 (SEQ ID NO: 8) cys17. 35 cysteine corresponds to that at human IL-171 ((SEQ ID NO: 3) cys50; at mouse IL-172 (SEQ ID NO: 12) cys141; at human IL-173

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(SEQ ID NO: 14) cys119; and mouse IL-174 (SEQ ID NO: 24) cys104. The disulfide linkages should be cysteines 2 with 5; and 3 with 6; and 1 with 4. Functional significance of the fold similarity suggests formation of dimers for the IL-170 family. As a consequence, IL-170 dimers would bring together two cell surface receptors, through which signal transduction will occur.

These new proteins are designated CTLA-8 related, or generally IL-170, proteins. The natural proteins should be capable of mediating various physiological responses which would lead to biological or physiological responses in target cells, e.g., those responses characteristic of cytokine signaling. Initial studies had localized the message encoding this protein to various cell lines of hematopoietic cells. Genes encoding the original CTLA-8 (IL-17) antigen have been mapped to mouse chromosome 1A and human chromosome 2q31.

Murine CTLA-8 was originally cloned by Rouvier, et al. (1993)

J. Immunol. 150:5445-5456. The human IL-173 has been mapped to chromosome 13q11. Similar sequences for proteins in other mammalian species should also be available.

Purified CTLA-8, when cultured with synoviocytes, is able to induce the secretion of IL-6 from these cells. This induction is reversed upon the addition of a neutralizing antibody raised against human CTLA-8. Endothelial, epithelial, fibroblast and carcinoma cells also exhibit responses to treatment with CTLA-8. This data suggests that CTLA-8 may be implicated in inflammatory fibrosis, e.g., psoriasis, sclerodermia, lung fibrosis, or cirrhosis. CTLA-8 may also cause proliferation of carcinomas or other cancer cells inasmuch as IL-6 often acts as a growth factor for such cells. As such, the newly discovered other related family members are likely to have similar or related biological activities.

The descriptions below are directed, for exemplary

35 purposes, to a murine or human IL-170 proteins, but are
likewise applicable to related embodiments from other species.

#### II. Nucleic Acids

Tables 1-6 disclose the nucleotide and amino acid sequences of various new IL-170 family member sequences. described nucleotide sequences and the related reagents are useful in constructing DNA clones useful for extending the clones in both directions for full length or flanking sequence detemination, expressing IL-170 polypeptides, or, e.g., isolating a homologous gene from another natural source. Typically, the sequences will be useful in isolating other genes, e.g., allelic variants, from mouse, and similar 10 procedures will be applied to isolate genes from other species, e.g., warm blooded animals, such as birds and mammals. Cross hybridization will allow isolation of genes from other species. A number of different approaches should be available to successfully isolate a suitable nucleic acid 15 clone from other sources.

20 Table 1: Nucleotide sequence encoding a primate, e.g., human, IL-171 under IUPAC code. Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 1:

25	GACACGGATG	AGGACCGCTA	TCCACAGAAG	CTGGCCTTCG	CCGAGTGCCT	GTGCAGAGGC	60
25	TGTATCGATG	CACGGACGGG	CCGCGAGACA	GCTGCGCTCA	ACTCCGTGCG	GCTGCTCCAG	120
	AGCCTGCTGG	TGCTGCGCCG	CCGGCCCTGC	TCCCGCGACG	GCTCGGGGCT	CCCCACACCT	180
30	GGGGCCTTTG	CCTTCCACAC	CGAGTTCATC	CACGTCCCCG	TCGGCTGCAC	CTGCGTGCTG	240
	CCCCGTTCAA	GTGTGACCGC	CAAGGCCGTG	GGGCCCTTAG	NTGACACCGT	GTGCTCCCCA	300 -
35	GAGGGACCCC	TATTTATGGG	AATTATGGTA	TTATATGCTT	CCCACATACT	TGGGGCTGGC	360
	ATCCCGNGCT	GAGACAGCCC	CCTGTTCTAT	TCAGCTATAT	GGGGAGAAGA	GTAGACTTTC	420
	AGCTAAGTGA	AAAGTGNAAC	GTGCTGACTG	TCTGCTGTCG	TNCTACTNAT	GCTAGCCCGA	480
40	GTGTTCACTC	TGAGCCTGTT	AAATATAGGC	GGTTATGTAC	С		521

SEQ ID NO: 2 and 3 are PATENTIN translatable cDNA and polypeptide sequences:

	Asp 1		Asp														40
50	CTG	TGC	AGA	GGC	TGT	ATC	GAT	GCA	CGG	ACG	GGC	CGC	GAG	ACA	GCT	GCG	96

CTG TGC AGA GGC TGT ATC GAT GCA CGG ACG GGC CGC GAG ACA GCT GCG
Leu Cys Arg Gly Cys Ile Asp Ala Arg Thr Gly Arg Glu Thr Ala Ala
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															CGC Arg			144
5															TTT Phe			192
10															GTG Val			240
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<b>2</b> 0															TTA Leu		٠	336
20								GGC Gly 120				TGA	GACA	GCC (	CCTC	STTCTA		389
25	TTC	AGCT	ATA 7	rggg	GAGAZ	AG AG	STAG2	ACTT	r cac	CTA	AGTG	AAA	AGTG1	ıAA (	CGTG	CTGACT		449
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55	cca Pro	cac His	tgc Cys	tac Tyr 20	tcg Ser	gct Ala	gag Glu	gaa Glu	ctg Leu 25	ccc Pro	ctc Leu	ggc Gly	cag Gln	gcc Ala 30	ccc Pro	cca Pro	261	٠
60															gta Val		309	
65															gag Glu		357	

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5															cgt Arg 95		453
10	gac Asp	acg Thr	gat Asp	gag Glu 100	gac Asp	cgc Arg	tat Tyr	cca Pro	cag Gln 105	aag Lys	ctg Leu	gcc Ala	ttc Phe	gcc Ala 110	gag Glu	tgc Cys	501
15	ctg Leu	tgc Cys	aga Arg 115	ggc Gly	tgt Cys	atc Ile	gat Asp	gca Ala 120	cgg Arg.	acg Thr	ggc Gly	cgc Arg	gag Glu 125	aca Thr	gct Ala	gcg Ala	549
20															cgc Arg		597
20	ccc Pro 145	tgc Cys	tcc Ser	cgc Arg	gac Asp	ggc Gly 150	tcg Ser	ggg Gly	ctc Leu	ccc Pro	aca Thr 155	cct Pro	ggg	gcc Ala	ttt Phe	gcc Ala 160	645
25	ttc Phe	cac His	acc Thr	gag Glu	ttc Phe 165	atc Ile	cac His	gtc Val	ccc Pro	gtc Val 170	ggc Gly	tgc Cys	acc Thr	tgc Cys	gtg Val 175	ctg Leu	693
30	ccc Pro				tgad	ccgcc	ga g	gccg	rtggg	gg co	ccta	agact	gga	acac	gtgt		745
	gctc	ccce	ıga ç	ggca	ccc	c ta	attta	atgto	, tat	ttat	tgg	tatt	tata	atg (	ected	cccaa	805
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	TGGC	CCAC	1					ieu I		CTG ? Leu !			Leu					109
<b>20</b>	TTT Phe	CTG Leu	AGT Ser -5	GAG Glu	GCG Ala	GCA Ala	GCT Ala	CGG Arg 1	AAA Lys	ATC Ile	CCC Pro	AAA Lys 5	GTA Val	GGA Gly	CAT His	ACT Thr		157
25 g	TTT Phe 10	TTC Phe	CAA Gln	AAG Lys	CCT Pro	GAG Glu 15	AGT Ser	TGC Cys	CCG Pro	CCT Pro	GTG Val 20	CCA Pro	GGA Gly	GGT Gly	AGT Ser	ATG Met 25		205
30 i	AAG Lys	CTT Leu	GAC Asp	ATT Ile	GGC Gly 30	ATC Ile	ATC Ile	AAT Asn	GAA Glu	AAC Asn 35	CAG Gln	CGC Arg	GTT Val	TCC Ser	ATG Met 40	TCA Ser		253
35	CGT Arg	AAC Asn	ATC Ile	GAG Glu 45	AGC Ser	CGC Arg	TCC Ser	ACC Thr	TCC Ser 50	CCC Pro	TGG Trp	AAT Asn	TAC Tyr	ACT Thr 55	GTC Val	ACT Thr		301
40	TGG Trp	GAC Asp	CCC Pro 60	Asn	CGG Arg	TAC Tyr	CCC Pro	TCG Ser 65	AAG Lys	TTG Leu	TAC Tyr	AGG Arg	CCC Pro 70	AAG Lys	TGT Cys	AGG Arg		349
1	AAC Asn	TTG Leu 75	GGC Gly	TGT Cys	ATC Ile	AAT Asn	GCT Ala 80	CAA Gln	GGA Gly	AAG Lys	GAA Glu	GAC Asp 85	ATC Ile	TnC Xaa	ATG Met	AAT Asn		397
	TCC Ser 90																	403

50
MVKYLLLSILGLAFLSEAAA RKIPKVGHTFFQKPESCPPVPGGSMKLDIGIINENQRVSMSRNIES
RSTSPWNYTVTWDPNRYPSKLYRPKCRNLGCINAQGKEDIXMNSV

Particularly interesting segments include, e.g., those which begin or end with arg1; cys17; pro18, pro19; val20; thr49; ser50; arg69; pro70; and the end of the sequence available.

60

5	Table 3: Nucleotide sequence encoding a primate, e.g., human, IL-172 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. Predicted signal cleavage site indicated, but may be a few residues on either side; putative glycosylation site at residues 55-57. SEQ ID NO: 9 and 10.
	at residues 55-57. SEQ ID NO: 9 and IU.

10				AAC Asn -15						48
15				CCC						96
20				CTG Leu						144
25				AAA Lys						192
				GTG Val 50						240
30				GTC Val						288
35				GJÀ GCC						336
40				GAG Glu						384
45				GAG Glu						432
				CGC Arg 130						480
50				CGC Arg						528
55	TGC Cys	 	TGA							543

<sup>60</sup> MDWPHNLLFLLTISIFLGLG QPRSPKSKRKGQGRPGPLVPGPHQVPLDLVSRMKPYARMEEYER NIEEMVAQLRNSSELAQRKCEVNLQLWMSNKRSLSPWGYSINHDPSRIPVDLPEARCLCLGCVNP FTMQEDRSMVSVPVFSQVPVRRRLCPPPPRTGPCRQRAVMETIAVGCTCIF

Particularly interesting segments include, e.g., those which begin or end with gln1; val19; pro20; pro22; lys34; pro35; leu78; ser79; glu98; ala99; phe110; thr111; cys143; or arg144.

5

10

Nucleotide sequence encoding a rodent, e.g., mouse, IL-172 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. Predicted signal cleavage site indicated, but may be a few residues on either side; putative glycosylation site at residues 53-55. SEQ ID NO: 11 and 12.

<b>1</b> 5						AGC Ser											48
20	CTG Leu	GCG Ala -5	CCA Pro	AGC Ser	CAC His	CCC Pro	CGG Arg 1	AAC Asn	ACC Thr	AAA Lys	GGC Gly 5	AAA Lys	AGA Arg	AAA Lys	GGG Gly	CAA Gln 10	96
25						TTG Leu											144
23						AAG Lys											192
. 30						GTG Val											240
35						GTC Val											288
40						GGC Gly 80											336
<b>4</b> 5						GAG Glu											384
						GAG Glu											432
50	AGC Ser	CAG Gļn	GTG Val 125	CCG Pro	GTG Val	CGC Arg	CGC Arg	CGC Arg 130	CTC Leu	TGT Cys	CCT Pro	CAA Gln	CCT Pro 135	CCT Pro	CGC Arg	CCT Pro	480
<b>55</b> .						CGT Arg											528
60	ACC Thr 155		ATC Ile		TGA												· 543

MDWPHSLLFLLAISIFLAPSHP RNTKGKRKGQGRPSPLAPGPHQVPLDLVSRVKPYARMEEYERN LGEMVAQLRNSSEPAKKKCEVNLQLWLSNKRSLSPWGYSINHDPSRIPADLPEARCLCLGCVNPFT MQEDRSMVSVPVFSQVPVRRRLCPQPPRPGPCRQRVVMETIAVGCTCIF

Particularly interesting segments include, e.g., those which begin or end with arg1; ala17; pro18; pro20; his21; lys32; pro33; leu76; ser77; glu96; ala97; phe108; thr109; cys141; or arg142.

٠5

Table 4: Nucleotide sequence encoding a primate, e.g., human, IL-173 polypeptide and predicted amino acid sequence.

10 Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 13 and 14.

15		GCG Ala															48
20	GCG Ala	GCC Ala	GGC Gly	GTG Val 20	CTC Leu	AGT Ser	GCC Ala	TTC Phe	CAC His 25	CAC His	ACG Thr	CTG Leu	CAG Gln	CTG Leu 30	GGG Gly	CCG Pro	96
25		GAG Glu															144
23		CGC Arg 50															192
30		TAC Tyr															<b>240</b>
35		GCC Ala															288
40	GAG Glu	GAC Asp	GTG Val	CGC Arg 100	TTC Phe	CGC Arg	AGC Ser	GCC Ala	CCT Pro- 105	GTC Val	TAC Tyr	ATG Met	CCC Pro	ACC Thr 110	GTC Val	GTC Val	336
45	CTG Leu	CGC Arg	CGC Arg 115	ACC Thr	CCC Pro	GCC Ala	TGC Cys	GCC Ala 120	GGC Gly	GGC Gly	CGT Arg	TCC Ser	GTC Val 125	TAC Tyr	ACC Thr	GAG Glu	384
45		TAC Tyr 130															432
50		GAC Asp						т									454

55

CADRPEELLEQLYGRLAAGVLSAFHHTLQLGPREQARNASCPAGGRPADRRFRTPTNLRS VSPWAYRISYDPARYPRYLPEAYCLCRGCLTGLFGEEDVRFRSAPVYMPTVVLRRTPACA GGRSVYTEAYVTIPVGCTCVPEPEKDADSIN Supplementary nucleotide sequence encoding a primate, e.g., human, IL-173 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. Predicted signal cleavage site indicated, but may be a few residues on either side. SEQ ID NO: 15 and 16.

	gcc	ggg	cag (	gtgg	cgac	ct c	gctca	agtc	g gct	tctc	ggt	ccaa	agtc	ccc (	gggto	tgg	58
10										ctg Leu							106
15										gcg Ala							154
20										ctg Leu 25							202
20										ctg Leu							250
25										ggg Gly							298
30										agc Ser							346
35	aga Arg 80	atc Ile	tcc Ser	tac Tyr	gac Asp	ccg Pro 85	gcg Ala	agg Arg	tac Tyr	ccc Pro	agg Arg 90	tac Tyr	ctg Leu	cct Pro	gaa Glu	gcc Ala 95	394.
40	tac Tyr	tgc Cys	ctg Leu	tgc Cys	cgg Arg 100	ggc Gly	tgc Cys	ctg Leu	acc Thr	ggg Gly 105	ctg Leu	ttc Phe	ggc Gly	gag Glu	gag Glu 110	gac Asp	442
40										atg Met							490
45										tcc Ser							538
50	gtc Val	acc Thr 145	atc Ile	ccc Pro	gtg Val	ggc Gly	tgc Cys 150	acc Thr	tgc Cys	gtc Val	ccc Pro	gag Glu 155	ccg Pro	gag Glu	aag Lys	gac Asp	586
55										aaa Lys							634
60			ccc Pro							ccc Pro 185	tgaç	ggcc	ggt (	ctg	cccg	gg	684
00	gagg	gtcto	ccc o	cggc	ccgca	at co	cgag	ggcgd	CC	aagct	gga	gcc	gcct	gga ç	gggct	cggtc	744
•	ggc	gacci	tct (	gaaga	agagt	tg ca	accga	agcaa	a acc	caagt	gcc	ggag	gcaco	cag d	egec	gccttt	804
65	ccat	tgga	gac 1	cgta	aagca	ag c	tcat	ctga	a cad	gggg	catc	cct	ggcti	tgc 1	cttta	agctac	864

	aagcaagcag cgtggctgga agctgatggg aaacgacccg gcacgggcat cctgtgtgcg 924
	gcccgcatgg agggtttgga aaagttcacg gaggctccct gaggagcctc tcagatcggc 984
5	tgctgcgggt gcagggcgtg actcaccgct gggtgcttgc caaagagata gggacgcata 1044
	tgctttttaa agcaatctaa aaataataat aagtatagcg actatatacc tacttttaaa 1104
10	atcaactgtt ttgaatagag gcagagctat tttatattat caaatgagag ctactctgtt 1164
	acatttetta acatataaac ategttttt acttettetg gtagaatttt ttaaageata 1224
	attggaatcc ttggataaat tttgtagctg gtacactctg gcctgggtct ctgaattcag 1284
15	cctgtcaccg atggctgact gatgaaatgg acacgtctca tctgacccac tcttccttcc 1344
	actgaaggtc ttcacgggcc tccaggcctc gtgccgaatt c 1385
20	MLVAGFLLALPPSWAAGAPRAGRRPARPRGCADRPEELLEQLYGRLAAGVLSAFHHTLQLGPREQARNAS CPAGGRPADRRFRPPTNLRSVSPWAYRISYDPARYPRYLPEAYCLCRGCLTGLFGEEDVRFRSAPVYMPT VVLRRTPACAGGRSVYTEAYVTIPVGCTCVPEPEKDADSINSSIDKQGAKLLLGPNDAPAGP
25	Important predicted motifs include, e.g., cAMP PK at 50-53, 66-69, 72-75, and 113-116; Ca Phos at 82-84 and 166-168; myristoly sites at 57-61 and 164-166; and phosphorylation sites at 50, 53, 72, 75, 80, 82, 113, and 116.
30	Nucleotide sequence encoding a rodent, e.g., rat, IL-173 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 17 and 18.
<b>3</b> 5	TTT CCG AGA TAC CTG CCC GAA GCC TAC TGC CTG TGC CGA GGC TGT CTG  Phe Pro Arg Tyr Leu Pro Glu Ala Tyr Cys Leu Cys Arg Gly Cys Leu  1 5 10 15
40	ACC GGG CTC TAC GGT GAG GAG GAC TTC CGC TTT CGC AGC GCA CCC GTC Thr Gly Leu Tyr Gly Glu Glu Asp Phe Arg Phe Arg Ser Ala Pro Val 20 25 30
<b>4</b> 5	TTC TCT CCG GCG GTG GTG CTG CGG CGC ACG GCC T  Phe Ser Pro Ala Val Val Leu Arg Arg Thr Ala Ala  35  40
	FPRYLPEAYCLCRGCLTGLYGEEDFRFRSAPVFSPAVVLRRTAA
50	Supplementary nucleotide sequence encoding a rodent, e.g., mouse, IL-173 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. Predicted signal cleavage site indicated, but may be a few residues on either side. SEQ ID NO: 19 and 20.
55	atg ttg ggg aca ctg gtc tgg atg ctc ctc gtc ggc ttc ctg ctg gca 48  Met Leu Gly Thr Leu Val Trp Met Leu Leu Val Gly Phe Leu Leu Ala -20 -15 -10
60	ctg gcg ccg ggc gcg gcg gcg ctg agg acc ggg agg cgc ccg 96 Leu Ala Pro Gly Arg Ala Ala Gly Ala Leu Arg Thr Gly Arg Arg Pro -5 -1 1 5

	gcg Ala	cgg Arg 10	ccg Pro	cgg Arg	gac Asp	tgc Cys	gcg Ala 15	gac Asp	cgg Arg	cca Pro	gag Glu	gag Glu 20	ctc Leu	ctg Leu	gag Glu	cag Gln	144
5	ctg Leu 25	tac Tyr	GJÀ aaa	cgg Arg	ctg Leu	gcg Ala 30	gcc Ala	ggc Gly	gtg Val	ctc Leu	agc Ser 35	gcc Ala	ttc Phe	cac His	cac His	acg Thr 40	192
10	ctg Leu	cag Gln	ctc Leu	ggg Gly	ccg Pro 45	cgc Arg	gag Glu	cag Gln	gcg Ala	cgc Arg 50	aat Asn	gcc Ala	agc Ser	tgc Cys	ccg Pro 55	gcc Ala	240
15	Gly ggg	ggc Gly	agg Arg	gcc Ala 60	gcc Ala	gac Asp	cgc Arg	cgc Arg	ttc Phe 65	cgg Arg	cca Pro	ccc Pro	acc Thr	aac Asn 70	ctg Leu	cgc Arg	288
20	agc Ser	gtg Val	tcg Ser 75	ccc Pro	tgg Trp	gcg Ala	tac Tyr	agg Arg 80	att Ile	tcc Ser	tac Tyr	gac Asp	cct Pro 85	gct Ala	cgc Arg	ttt Phe	336
								tac Tyr									384
25	Gly 105	Leu	Tyr	Gly	Glu	Glu 110	Asp	ttc Phe	Arg	Phe	Arg 115	Ser	Thr	Pro	Val	Phe 120	432
30	tct Ser	cca Pro	gcc Ala	gtg Val	gtg Val 125	ctg Leu	cgg Arg	cgc Arg	aca Thr	gcg Ala 130	gcc Ala	tgc Cys	gcg Ala	ggc Gly	ggc Gly 135	ege Arg	480
25	tct Ser	gtg Val	tac Tyr	gcc Ala 140	gaa Glu	cac His	tac Tyr	atc Ile	acc Thr 145	atc Ile	ccg Pro	gtg Val	ggc Gly	tgc Cys 150	acc Thr	tgc Cys	528
35	gtg Val	ccc Pro	gag Glu 155	ccg Pro	gac Asp	aag Lys	tcc Ser	gcg Ala 160	gac Asp	agt Ser	gcg Ala	aac Asn	tcc Ser 165	agc Ser	atg Met	gac Asp	576
40	aag Lys	ctg Leu 170	ctg Leu	ctg Leu	ggg Gly	ccc Pro	gcc Ala 175	gac Asp	agg Arg	cct Pro	gcg Ala	ggg Gly 180	cgc Arg	tgat	geeç	<b>3</b> gg	625
45	gact	gcco	ege o	atgg	gccca	ag ct	tcct	gcat	gca	tcag	gtc	ccct	ggcc	ct o	gacaa	aaaccc	685
40	acco	cato	gat d	ccto	gccg	jc to	gccta	attt	tto	caaa	agg	acag	ctac	at a	agct	ttaaa	745
•	tata	attt	ttc a	aaagt	agad	ca ct	acat	atct	: aca	acta	ttt	tgaa	tagt	gg	cagaa	actat	805
50				-		_	_	_	_							aagcac	
																ctcaga	
55																gcttac	
										•						agaggg	
60												CCCa	igctt	at (	gato	ggtctt	
60 .	aact	ttai	taa a	agatt	caaag	jc ct	ttgg	gtgtt	. att	CTTT	.c						1143

MLGTLVWMLLVGFLLALAPGRAAGALRTGRRPARPRDCADRPEELLEQLYGRLAAGVLSAFHHTL QLGPREQARNASCPAGGRAADRRFRPPTNLRSVSPWAYRISYDPARFPRYLPEAYCLCRGCLTGL YGEEDFRFRSTPVFSPAVVLRRTAACAGGRSVYAEHYITIPVGCTCVPEPDKSADSANSSMDKLL LGPADRPAGR.

55

60

Important predicted motifs include, e.g., cAMP PK sites at 50-53, 66-69, 72-75, and 113-116; Ca phosphorylation sites at 82-84, 159-161, and 166-168; myristoly sites at 57-61 and 101-105; N-glycosyl sites at 51-53 and 164-166; phosphorylation sites at 50, 53, 72, 75, 80, 82, 113, and 116; and PKC phosphorylation sites at 4-6

10 Table 5: Nucleotide sequence encoding a primate, e.g., human, IL-174 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 21 and 22.

tgagtgtgca gtgccagc atg tac cag gtg gtt gca ttc ttg gca atg gtc	51
Met Tyr Gln Val Val Ala Phe Leu Ala Met Val	
-15 -10	

- 20 atg gga acc cac acc tac agc cac tgg ccc agc tgc tgc ccc agc aaa 99
  Met Gly Thr His Thr Tyr Ser His Trp Pro Ser Cys Cys Pro Ser Lys
  -5 -1 1 5 10
- ggg cag gac acc tct gag gag ctg ctg agg tgg agc act gtg cct gtg 147
  25 Gly Gln Asp Thr Ser Glu Glu Leu Leu Arg Trp Ser Thr Val Pro Val
  15 20 25
- cct ccc cta gag cct gct agg ccc aac cgc cac cca gag tcc tgt agg 195
  Pro Pro Leu Glu Pro Ala Arg Pro Asn Arg His Pro Glu Ser Cys Arg
  30 35 40
  - gcc agt gaa gat gga ccc ctc aac agc agg gcc atc tcc ccc tgg aga 243 Ala Ser Glu Asp Gly Pro Leu Asn Ser Arg Ala Ile Ser Pro Trp Arg 45 55
    - tat gag ttg gac aga gac ttg aac cgg ctc ccc cag gac ctg tac cac
      Tyr Glu Leu Asp Arg Asp Leu Asn Arg Leu Pro Gln Asp Leu Tyr His
      60 65 70 75
- 40 gcc cgt tgc ctg tgc ccg cac tgc gtc agc cta cag aca ggc tcc cac 339
  Ala Arg Cys Leu Cys Pro His Cys Val Ser Leu Gln Thr Gly Ser His
  80 85 90
- atg gac ccc cgg ggc aac tcg gag ctg ctc tac cac aac cag act gtc 387
  45 Met Asp Pro Arg Gly Asn Ser Glu Leu Leu Tyr His Asn Gln Thr Val
  95 100 105
- ttc tac cgg cgg cca tgc cat ggc gag aag ggc acc cac aag ggc tac

  Phe Tyr Arg Arg Pro Cys His Gly Glu Lys Gly Thr His Lys Gly Tyr

  110

  115

  435
  - tgc ctg gag cgc agg ctg tac cgt gtt tcc tta gct tgt gtg tgt gtg

    Cys Leu Glu Arg Arg Leu Tyr Arg Val Ser Leu Ala Cys Val Cys Val

    125

    130

    483

cgg ccc cgt gtg atg ggc tag 504
Arg Pro Arg Val Met Gly
140 145

MYQVVAFLAMVMGTHTYSHWPSCCPSKGQDTSEELLRWSTVPVPPLEPARPNRHPESCRASED GPLNSRAISPWRYELDRDLNRLPQDLYHARCLCPHCVSLQTGSHMDPRGNSELLYHNQTVFYR RPCHGEKGTHKGYCLERRLYRVSLACVCVRPRVMG

Important predicted motifs include, e.g., cAMP PK sites at 21-24, 53-56, and 95-98; Ca phosphorylation sites at 15-17, 16-18, and 45-47; myristoly sites at 12-16, 115-119, and 118-122; N-glycosyl site at 104-107; phosphorylation sites at 21, 23, 43, 53, 56, 95, 98, and 131; PKC phosphorylation sites at 41-43 and 119-121; and tyrosine kinase site at 95-102. 10 Nucleotide sequence encoding a rodent, e.g., mouse, IL-174 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. ID NO: 23 and 24. 15 CGG CAC AGG CGG CAC AAA GCC CGG AGA GTG GCT GAA GTG GAG CTC TGC 48 Arg His Arg Arg His Lys Ala Arg Arg Val Ala Glu Val Glu Leu Cys 10 ATC TGT ATC CCC CCC AGA GCC TCT GAG CCA CAC CCA CGC AGA ATC 96 20 Ile Cys Ile Pro Pro Arg Ala Ser Glu Pro His Pro Pro Arg Arg Ile CTG CAG GGC CAG CAA GGA TGG CCT CTC AAC AGC AGG GCC ATC TCT CCT 144 Leu Gln Gly Gln Gln Gly Trp Pro Leu Asn Ser Arg Ala Ile Ser Pro 25 35 40 TGG AGC TAT GAG TTG GAC AGG GAC TTG AAT CGG GTC CCC CAG GAC TGG 192 Trp Ser Tyr Glu Leu Asp Arg Asp Leu Asn Arg Val Pro Gln Asp Trp 30 TAC CAC GCT CGA TGC CTG TGC CCA CAC TGC GTC ACG CTA CAG ACA GGC 240 Tyr His Ala Arg Cys Leu Cys Pro His Cys Val Thr Leu Gln Thr Gly 35 TCC CAC ATG GAC CCG CTG GGC AAC TCC GTC CCA CTT TAC CAC AAC CAG 288 Ser His Met Asp Pro Leu Gly Asn Ser Val Pro Leu Tyr His Asn Gln ACG GTC TTC TAC CGG CGG CCA TGC ATG GCG AGG AAG GTA CCC ATC GCC 336 40 Thr Val Phe Tyr Arg Arg Pro Cys Met Ala Arg Lys Val Pro Ile Ala 100 105 GCT ACT GCT TGG AGC GCA GGT CTA CCG AGT CTC CTT GGC TTG TGT GTG 384 Ala Thr Ala Trp Ser Ala Gly Leu Pro Ser Leu Leu Gly Leu Cys Val 45 120 TGT GCG GCC CCG GGT CAT GGC TTA GTC ATG CTC ACC ATC TGC CTG AGG 432 Cys Ala Ala Pro Gly His Gly Leu Val Met Leu Thr Ile Cys Leu Arg 130 50 TGAATGCCGG GTGGGAGAGA GGGCCAGGTG TACATCACCT GCCAATGCGG GCCGGGTTCA 492 AGCCTGCAAA GCCTACCTGA AGCAGCAGGT CCCGGGACAG GATGGAGACT TGGGGAGAAA 552 55 TCTGACTTTT GCACTTTTTG GAGCATTTTG GGAAGAGCAG GTTCGCTTGT GCTGTAGAGA 612 TGCTGTTG 620

60 RHRRHKARRVAEVELCICIPPRASEPHPPRRILQGQQGWPLNSRAISPWSYELDRDLNRVPQDWYHARC LCPHCVTLQTGSHMDPLGNSVPLYHNQTVFYRRPCMARKVPIAATAWSAGLPSLLGLCVCAAPGHGLVM LTICLR

Supplementary nucleotide sequence encoding a rodent, e.g., mouse, IL-174 polypeptide and predicted amino acid sequence. Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 25 and 26.

5	pur	pos	es.	SEQ ID NO: 25 and 26.													
								ttg Leu									48
10								ggc Gly									96
15	ccc	agc Ser	aaa Lys	gag Glu 20	Gln	gaa Glu	ccc Pro	ccg Pro	gag Glu 25	gag Glu	tgg Trp	ctg Leu	aag Lys	tgg Trp 30	agc Ser	tct Ser	144
20								cct Pro 40									192
20								ggc Gly									240
25								agg Arg									288
30	ctg Leu							tgc Cys									336
35	ggc Gly	tcc Ser	cac	atg Met 100	gac Asp	ccg Pro	ctg Leu	ggc Gly	aac Asn 105	tcc Ser	gtc Val	cca Pro	ctt Leu	tac Tyr 110	cac His	aac Asn	384
40	cag Gln	acg Thr	gtc Val 115	ttc Phe	tac Tyr	cgg Arg	cgg Arg	cca Pro 120	tgc Cys	cat His	ggt Gly	gag Glu	gaa Glu 125	ggt Gly	acc Thr	cat His	432
40								agg Arg									480
45								atg Met		tagt	catg	et c	acca	cctg	ic .		527
50	ctga	ıggct	ga t	gccc	ggtt	g gg	gagag	aggg	cca	ggtg	rtac	aato	acct	tg d	caat	gcggg	587
30	ccgg	gtto	caa g	jccct	ccaa	a go	ccta	cctg	aag	cago	agg	ctcc	cggg	ac a	agat	ggagg	647
	actt	gggg	gag a	aact	ctga	c tt	ttgc	actt	ttt	.ggaa	gca	cttt	tggg	aa g	gago	aggtt	707
55	ccgc	ttgt	gc t	gcta	ıgagç	ra to	gctgt	tgtg	gca	tttc	tac	tcag	gaac	gg a	ctcc	aaagg	767
	cctg	ctga	cc c	tgga	agco	a ta	ctcc	tggc	tcc	tttc	ccc	tgaa	tccc	cc a	acto	ctggc	827
60	acag	gcac	tt t	ctcc	acct	c to	cccc	tttg	cct	tttg	ttg	tgtt	tgtt	tg t	gcat	gccaa	887
	ctct	gcgt	gc a	igcca	ggtg	rt aa	ttgc	cttg	aag	gatg	gtt	ctga	ıggtg	raa a	gctg	ttatc	947
	gaaa	igtga	ag a	ıgatt	tato	c aa	ataa	acat	ctg	tgtt	t						985

MYQAVAFLAMIVGTHTVSLRIQEGCSHLPSCCPSKEQEPPEEWLKWSSASVSPPEPLSHTHHAESCRAS KDGPLNSRAISPWSYELDRDLNRVPQDLYHARCLCPHCVSLQTGSHMDPLGNSVPLYHNQTVFYRRPCH GEEGTHRRYCLERRLYRVSLACVCVRPRVMA

Important predicted motifs include, e.g., cAMP PK sites at 29-32 and 61-64; Ca phosphorylation sites at 18-20, 53-55, and 67-69; myristoly site at 123-127; N-glycosylation site at 112-114; and phosphorylation sites at 29, 31, 51, 53, 61, 64, 139, and 141; and PKC phosphorylation sites at 2-4, 49-51, and 127-129.

- Table 6: Nucleotide sequence encoding a primate, e.g., human, 15 IL-176. Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 27 and 28:
- tat tac ttg tta ggg aga ccc aat ggt agt ttt att cct tgg gga tac 95
  Tyr Tyr Leu Leu Gly Arg Pro Asn Gly Ser Phe Ile Pro Trp Gly Tyr
  25 20 25 30
  - ata gta aat act tca tta aag tcg agt aca gaa ttt gat gaa aag tgt 143
    Ile Val Asn Thr Ser Leu Lys Ser Ser Thr Glu Phe Asp Glu Lys Cys
    35 40 45
  - gga tgt gtg gga tgt act gcc gcc ttc aga agt cca cac act gcc tgg 191 Gly Cys Val Gly Cys Thr Ala Ala Phe Arg Ser Pro His Thr Ala Trp 50 55 60
- 35 agg gag aga act gct gtt tat tca ctg att aag cat ttg ctg tgt acc 239
  Arg Glu Arg Thr Ala Val Tyr Ser Leu Ile Lys His Leu Leu Cys Thr
  65 70 75
- aac tac ttt tca tgt ctt atc tta att ctc ata aca gtc att
  40 Asn Tyr Phe Ser Cys Leu Ile Leu Ile Leu Ile Thr Val Ile
  80 85 90
  - tgatatttta aaaaacccca gaaatctgag aaagagataa agtggtttgc tcaaggttat 341
- 45 agaacagact accatgtgtt gtatttcaga ttttaattca tgtttgtctg attttaagtt 401
  - ttgttcgctt gccagggtac cccacaaaaa tgccaggcag ggcattttca tgatgcactt 461
- gagatacctg aaatgacagg gtagcatcac acctgagagg ggtaaaggat gggaacctac 521
  - cttccatggc cgctgcttgg cagtctcttg ctgcatgcta gcagagccac tgtatatgtg 581
  - ccgaggctct gagaattaac tgcttaaaga actgccttct ggagggagaa gagcacaaga 641
- 55 tcacaattaa ccatatacac atcttactgt gcgaggtcat tgagcaatac aggagggatt 701
- ttatacattt tagcaactat cttcaaaacc tgagctatag ttgtattctg cccccttcct 761

ctgggcaaaa gtgtaaaagt ttg 784

VPYLFKKIILHFFASYYLLGRPNGSFIPWGYIVNTSLKSSTEFDEKCGCVGCTAAFRSPHTAWRERTAVYS LIKHLLCTNYFSCLILILITVI

5	Nucleotide sequence encoding a primate, e.g., human, IL-177. Also can use complementary nucleic acid sequences for many purposes. SEQ ID NO: 29 and 30:										
	gtg act gta ttg tgg gga cag gaa gca caa att ccc atg tgg atc act 48 Val Thr Val Leu Trp Gly Gln Glu Ala Gln Ile Pro Met Trp Ile Thr 1 5 10 15										
10	agg aga gat aat aag tgg ggt cat ttc acc cct tgg tcc cct gct tcc 96 Arg Arg Asp Asn Lys Trp Gly His Phe Thr Pro Trp Ser Pro Ala Ser 20 25 30										
15	aga ccc aaa gag gcc tac atg gca ttg tgc ttc ctt ctt agt tgt agg 144 Arg Pro Lys Glu Ala Tyr Met Ala Leu Cys Phe Leu Leu Ser Cys Arg 35 40 45										
20	agg tgt gag ata caa tca ttt gcc tct gac ttt gag ggt tgg tcc 189 Arg Cys Glu Ile Gln Ser Phe Ala Ser Asp Phe Glu Gly Trp Ser 50 55 60										
	tagcatgccc ctgaccagta gccccttaaa tacttcattg atatggaagg tctctgaatc 249										
25	ttcgtgggct taatctacca ctctctgaag ttcttatgtc tttcaaaggc ctctaaaatc 309										
	tetgecatgt ettgeteate eagttgttag eatgatgtea ttgatacagt ggaetttgga 369 ~										
	atctaagtgg ggagacactg gtaagtgacc aattacttca cctgtggtgt gcaagccaga 429										
30	tcaggaagcc tctacctgca cgacaacaca t 460										
	VTVLWGQEAQIPMWITRRDNKWGHFTPWSPASRPKEAYMALCFLLSCRRCEIQSFASDFEGWS										
<b>35</b>											
40	Table 7: Alignment of various CTLA-8/IL-170 family members. The rat CTLA-8 sequence is SEQ ID NO: 31 (see GB L13839; 293329/30); mouse CTLA-8 sequence is SEQ ID NO: 32 (see GB 1469917/8); human CTLA-8 is SEQ ID NO: 33 (see GB U32659; 115222/3); and Herpes Saimiri virus ORF13 is SEQ ID NO: 34 (see GB Y13183; 2370235). CLUSTAL X (1.64b) multiple sequence alignment										
45											
50	IL-74_MuMYQAVAFLAMIVGTHTVSLRIQEGCSHLPSCCPSKEQEPPEEWLKWS IL-74_HuMYQVVAFLAMVMGTHTYSHWPSCCPSKEQDTSEELLRWS IL-72_HuMDWPHNLLFILLTISIFLGLGQPRSPKSKRKGQGRPGPLVPGPHQVPLDLVSRMK IL-72_MuMDWPHSLLFLLAISIFLAPSHPRNTKGKRKGQGRPSPLAPGPPQVPLDLVSRVK IL-73_Mu										
55	IL-17_RtMCLMLLLLLNLEATVKAAVLIPQSSVCPNAEANNFLQNVKVNL IL-17_MuMLLLLLSLAATVKAAAIIPQSSACPNTEAKDFLQNVKVNL IL-75_HuMVKYLLLSILGLAFLSEAAARKIPKVGHTFFQKPESCPPVPGGSMKLDIGIIN IL-71_Hu MTLLPGLLFLTWLHTCLAHHDPSLRGHPHSHGTPHCYSAEELPLGQAPPHLLARGAKWGQ										

5	IL-74_Mu IL-74_Hu IL-72_Hu IL-72_Mu IL-73_Mu IL-73_Hu IL-17_Hu IL-17_Hs IL-17_Mu IL-17_Mu IL-75_Hu IL-75_Hu	SASVSPP-EPLSHTHHAESCRASKD-GPLNSRAISPWSYELDRDLNRV TVPVPPL-EPARPNRHPESCRASED-GPLNSRAISPWRYELDRDLNRL P-YARMEEYERNIEEMVAQLRNSSELAQ-RKCEVNLQLWMSNKRSLSPWGYSINHDPSRI P-YARMEEYERNLGEMVAQLRNSSEPAK-KKCEVNLQLWLSNKRSLSPWGYSINHDPSRI AGVLSAFHHTLQLGPR-EQARNASCPAGGRAADRRFR-PPTNLRSVSPWAYRISYDPARF AGVLSAFHHTLQLGPR-EQARNASCPAGGRPADRRFR-PPTNLRSVSPWAYRISYDPARY NIHNRNTNTNP-KRSSDYYNRSTSPWNLHRNEDPERY SIRNWNTSSKRASDYYNRSTSPWTLHRNEDPDRY KVINSLSKASSRRPSDYLNRSTSPWTLHRNEDPDRY KVINSLGAKVSSRRPSDYLNRSTSPWTLHRNEDPDRY EN-QRVSMSR-NIESRSTSPWNYTVTWDPNRY ALPVALVSSLEAASHRGRHERPSATTQCPVLRPEEVLEADTHQRSISPWRYRVDTDEDRY *: ***
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20 25	IL-74_Mu IL-74_Hu IL-72_Hu IL-72_Mu IL-73_Mu IL-73_Hu IL-17_Hu IL-17_Hs IL-17_Rt IL-17_Mu IL-75_Hu IL-75_Hu IL-75_Hu	PQDLYHARCLCPHCVSLQTGSHMDPLGNSVPLYHNQTVFYRRPCHGEEGTHRRYCLER PQDLYHARCLCPHCVSLQTGSHMDPRGNSELLYHNQTVFYRRPCHGEKGTHKGYCLER PVDLPEARCLCLGCVNPFTM-QEDRSMVSVPVFS-QVPVRRRLCPPPPRTGPCRQR PADLPEARCLCLGCVNPFTM-QEDRSMVSVPVFS-QVPVRRRLCPQPPRPGPCRQR PRYLPEAYCLCRGCLTGLYG-EEDFRFRSTPVFS-PAVVLRRTAACAGGRSVYA PRYLPEAYCLCRGCLTGLFG-EEDVRFRSAPVYM-PTVVLRRTPACAGGRSVYT PSVIWEAKCRHLGCINADGNVDYHMNSVPIQQEILVLRREPPHCPNSFR PSVIWEAKCRYLGCVNADGNVDYHMNSVPIQQEILVVRKGHQPCPNSFR PSVIWEAQCRHQRCVNAEGKLDHHMNSVLIQQEILVLKREPEKCPFTFR PSVIWEAQCRHQRCVNAEGKLDHHMNSVLIQQEILVLKREPESCPFTFR PSEVVQAQCRNLGCINAQGKEDISMNSVPIQQETLVVRRKHQGCSVSFQ PQKLAFAECLCRGCIDARTG-RETAALNSVRLLQSLLVLRRRPCSRDGSGLPTPGAFAFH
30		• " • • • • • • • • • • • • • • • • • •
35 40	IL-74_Mu IL-74_Hu IL-72_Hu IL-72_Mu IL-73_Mu IL-73_Hu IL-17_Hu IL-17_Hs IL-17_Rt IL-17_Mu IL-75_Hu	RLYR-VSLACVCVRPRVMA
	IL-71_Hu	TEFIHVPVGCTCVLPRSV

Particularly interesting segments include, e.g., those corresponding to the segments of IL-172 or IL-175, indicated above, with the other family members.

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Purified protein or polypeptides are useful for generating antibodies by standard methods, as described above. Synthetic peptides or purified protein can be presented to an immune system to generate a specific binding composition, e.g., monoclonal or polyclonal antibodies. See, e.g., Coligan (1991) Current Protocols in Immunology Wiley/Greene; and Harlow and Lane (1989) Antibodies: A Laboratory Manual Cold Spring Harbor Press.

For example, the specific binding composition could be used for screening of an expression library made from a cell line which expresses an IL-170 protein. The screening can be standard staining of surface expressed protein, or by panning. Screening of intracellular expression can also be performed by various staining or immunofluorescence procedures. The binding compositions could be used to affinity purify or sort out cells expressing the protein.

This invention contemplates use of isolated DNA or fragments to encode a biologically active corresponding IL-170 protein or polypeptide. In addition, this invention covers isolated or recombinant DNA which encodes a biologically active protein or polypeptide and which is capable of hybridizing under appropriate conditions with the DNA sequences described herein. Said biologically active protein or polypeptide can be an intact antigen, or fragment, and have an amino acid sequence as disclosed in Tables 1-6. Further, this invention covers the use of isolated or recombinant DNA, or fragments thereof, which encode proteins which are homologous to an IL-170 protein or which were isolated using cDNA encoding an IL-170 protein as a probe. The isolated DNA can have the respective regulatory sequences in the 5' and 3' flanks, e.g., promoters, enhancers, poly-A addition signals, and others.

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An "isolated" nucleic acid is a nucleic acid, e.g., an RNA, DNA, or a mixed polymer, which is substantially separated from other components which naturally accompany a native sequence, e.g., ribosomes, polymerases, and flanking genomic sequences from the originating species. The term embraces a nucleic acid sequence which has been removed from its naturally occurring environment, and includes recombinant or cloned DNA isolates and chemically synthesized analogs or analogs biologically synthesized by heterologous systems. A substantially pure molecule includes isolated forms of the molecule. Alternatively, a purified species may be separated from host components from a recombinant expression system.

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The size of homology of such a nucleic acid will typically be less than large vectors, e.g., less than tens of kB, typically less than several kB, and preferably in the 2-6 kB range.

An isolated nucleic acid will generally be a homogeneous composition of molecules, but will, in some embodiments, contain minor heterogeneity. This heterogeneity is typically found at the polymer ends or portions not critical to a desired biological function or activity.

A "recombinant" nucleic acid is defined either by its method of production or its structure. In reference to its method of production, e.g., a product made by a process, the process is use of recombinant nucleic acid techniques, e.g., involving human intervention in the nucleotide sequence, typically selection or production. Alternatively, it can be a nucleic acid made by generating a sequence comprising fusion of two fragments which are not naturally contiguous to each other, but is meant to exclude products of nature, e.g., naturally occurring mutants. Thus, for example, products made by transforming cells with any unnaturally occurring vector is encompassed, as are nucleic acids comprising sequence derived using any synthetic oligonucleotide process. Such is often done to replace a codon with a redundant codon encoding the same or a conservative amino acid, while typically introducing or removing a sequence recognition site. Alternatively, it is performed to join together nucleic acid segments of desired functions to generate a single genetic entity comprising a desired combination of functions not found in the commonly available natural forms. Restriction enzyme recognition sites are often the target of such artificial manipulations, but other site specific targets, e.g., promoters, DNA replication sites, regulation sequences, control sequences, or other useful features may be incorporated by design. A similar concept is intended for a recombinant, e.g., fusion, polypeptide. Specifically included are synthetic nucleic acids which, by genetic code redundancy, encode polypeptides similar to fragments of these antigens, and fusions of sequences from various different species variants.

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A significant "fragment" in a nucleic acid context is a contiguous segment of at least about 17 nucleotides, generally at least 20 nucleotides, more generally at least 23 nucleotides, ordinarily at least 26 nucleotides, more ordinarily at least 29 nucleotides, often at least 32 nucleotides, more often at least 35 nucleotides, typically at least 38 nucleotides, more typically at least 41 nucleotides, usually at least 44 nucleotides, more usually at least 47 nucleotides, preferably at least 50 nucleotides, more preferably at least 53 nucleotides, and in particularly preferred embodiments will be at least 56 or more nucleotides. Said fragments may have termini at any location, but especially at boundaries between structural domains.

In other embodiments, the invention provides polynucleotides (or polypeptides) which comprise a plurality of distinct, e.g., nonoverlapping, segments of the specified length. Typically, the plurality will be at least two, more usually at least three, and preferably 5, 7, or even more. While the length minima are provided, longer lengths, of 20 various sizes, may be appropriate, e.g., one of length 7, and two of length 12.

A DNA which codes for an IL-170 protein will be particularly useful to identify genes, mRNA, and cDNA species which code for related or homologous proteins, as well as DNAs which code for homologous proteins from different species. There are likely homologues in other species, including primates. Various CTLA-8 proteins should be homologous and are encompassed herein. However, even proteins that have a more distant evolutionary relationship to the antigen can readily be isolated under appropriate conditions using these sequences if they are sufficiently homologous. Primate CTLA-8 protein proteins are of particular interest.

This invention further covers recombinant DNA molecules and fragments having a DNA sequence identical to or highly 35 homologous to the isolated DNAs set forth herein. particular, the sequences will often be operably linked to DNA segments which control transcription, translation, and DNA

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replication. Alternatively, recombinant clones derived from the genomic sequences, e.g., containing introns, will be useful for transgenic studies, including, e.g., transgenic cells and organisms, and for gene therapy. See, e.g., Goodnow (1992) "Transgenic Animals" in Roitt (ed.) Encyclopedia of Immunology Academic Press, San Diego, pp. 1502-1504; Travis (1992) Science 256:1392-1394; Kuhn, et al. (1991) Science 254:707-710; Capecchi (1989) Science 244:1288; Robertson (ed. 1987) Teratocarcinomas and Embryonic Stem Cells: A Practical Approach IRL Press, Oxford; Rosenberg (1992) J. Clinical Oncology 10:180-199; and Cournoyer and Caskey (1993) Ann. Rev. Immunol. 11:297-329.

Homologous nucleic acid sequences, when compared, exhibit significant similarity. The standards for homology in nucleic acids are either measures for homology generally used in the art by sequence comparison or based upon hybridization conditions. The hybridization conditions are described in greater detail below.

Substantial homology in the nucleic acid sequence comparison context means either that the segments, or their 20 complementary strands, when compared, are identical when optimally aligned, with appropriate nucleotide insertions or deletions, in at least about 50% of the nucleotides, generally at least 56%, more generally at least 59%, ordinarily at least 62%, more ordinarily at least 65%, often at least 68%, more 25 often at least 71%, typically at least 74%, more typically at least 77%, usually at least 80%, more usually at least about 85%, preferably at least about 90%, more preferably at least about 95 to 98% or more, and in particular embodiments, as high at about 99% or more of the nucleotides. Alternatively, 30 substantial homology exists when the segments will hybridize under selective hybridization conditions, to a strand, or its complement, typically using a sequence derived from Table 1, 2, or 3. Typically, selective hybridization will occur when there is at least about 55% homology over a stretch of at 35 least about 14 nucleotides, preferably at least about 65%, more preferably at least about 75%, and most preferably at

least about 90%. See, Kanehisa (1984) <u>Nuc. Acids Res.</u> 12:203-213. The length of homology comparison, as described, may be over longer stretches, and in certain embodiments will be over a stretch of at least about 17 nucleotides, usually at least about 20 nucleotides, more usually at least about 24 nucleotides, typically at least about 28 nucleotides, more typically at least about 40 nucleotides, preferably at least about 50 nucleotides, and more preferably at least about 75 to 100 or more nucleotides.

10 Stringent conditions, in referring to homology in the hybridization context, will be stringent combined conditions of salt, temperature, organic solvents, and other parameters, typically those controlled in hybridization reactions. Stringent temperature conditions will usually include 15 temperatures in excess of about 30° C, more usually in excess of about 37° C, typically in excess of about 45° C, more typically in excess of about 55° C, preferably in excess of about 65° C, and more preferably in excess of about 70° C. Stringent salt conditions will ordinarily be less than about 20 1000 mM, usually less than about 500 mM, more usually less than about 400 mM, typically less than about 300 mM, preferably less than about 200 mM, and more preferably less than about 150 mM. However, the combination of parameters is much more important than the measure of any single parameter. See, e.g., Wetmur and Davidson (1968) J. Mol. Biol. 31:349-25 370. Hybridization under stringent conditions should give a background of at least 2-fold over background, preferably at least 3-5 or more.

Alternatively, for sequence comparison, typically one
sequence acts as a reference sequence, to which test sequences
are compared. When using a sequence comparison algorithm,
test and reference sequences are input into a computer,
subsequence coordinates are designated, if necessary, and
sequence algorithm program parameters are designated. The
sequence comparison algorithm then calculates the percent
sequence identity for the test sequence(s) relative to the

reference sequence, based on the designated program parameters.

Optical alignment of sequences for comparison can be conducted, e.g., by the local homology algorithm of Smith and Waterman (1981) <a href="Adv. Appl. Math.">Adv. Appl. Math.</a> 2:482, by the homology alignment algorithm of Needleman and Wunsch (1970) <a href="J. Mol. Biol.">J. Mol.</a> Biol. 48:443, by the search for similarity method of Pearson and Lipman (1988) <a href="Proc. Nat'l Acad. Sci. USA">Proc. Nat'l Acad. Sci. USA</a> 85:2444, by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by visual inspection (see generally Ausubel, et al., supra).

One example of a useful algorithm is PILEUP. PILEUP 15 creates a multiple sequence alignment from a group of related sequences using progressive, pairwise alignments to show relationship and percent sequence identity. It also plots a tree or dendogram showing the clustering relationships used to create the alignment. PILEUP uses a simplification of the progressive alignment method of Feng and Doolittle (1987) J. 20 Mol. Evol. 35:351-360. The method used is similar to the method described by Higgins and Sharp (1989) CABIOS 5:151-153. The program can align up to 300 sequences, each of a maximum length of 5,000 nucleotides or amino acids. The multiple alignment procedure begins with the pairwise alignment of the 25 two most similar sequences, producing a cluster of two aligned sequences. This cluster is then aligned to the next most related sequence or cluster of aligned sequences. clusters of sequences are aligned by a simple extension of the pairwise alignment of two individual sequences. The final 30 alignment is achieved by a series of progressive, pairwise alignments. The program is run by designating specific sequences and their amino acid or nucleotide coordinates for regions of sequence comparison and by designating the program 35 parameters. For example, a reference sequence can be compared to other test sequences to determine the percent sequence identity relationship using the following parameters: default

gap weight (3.00), default gap length weight (0.10), and weighted end gaps.

Another example of algorithm that is suitable for determining percent sequence identity and sequence similarity 5 is the BLAST algorithm, which is described Altschul, et al. (1990) <u>J. Mol. Biol.</u> 215:403-410. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http:www.ncbi.nlm.nih.gov/). This algorithm involves first 10 identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul, et al., supra). These initial neighborhood word 15 hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are then extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Extension of the word hits 20 in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The 25 BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLAST program uses as defaults a wordlength (W) of 11, the BLOSUM62 scoring matrix (see Henikoff and Henikoff (1989) Proc. Nat'l Acad. Sci. USA 89:10915) alignments (B) of 50, expectation (E) of 30 10, M=5, N=4, and a comparison of both strands.

In addition to calculating percent sequence identity, the BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin and Altschul (1993) Proc. Nat'l Acad. Sci. USA 90:5873-5787). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or

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amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.1, more preferably less than about 0.01, and most preferably less than about 0.001.

A further indication that two nucleic acid sequences of polypeptides are substantially identical is that the polypeptide encoded by the first nucleic acid is immunologically cross reactive with the polypeptide encoded by the second nucleic acid, as described below. Thus, a polypeptide is typically substantially identical to a second polypeptide, for example, where the two peptides differ only by conservative substitutions. Another indication that two nucleic acid sequences are substantially identical is that the two molecules hybridize to each other under stringent conditions, as described below.

CTLA-8-like proteins from other mammalian species can be cloned and isolated by cross-species hybridization of closely related species, e.g., human, as disclosed in Tables 1-5. Homology may be relatively low between distantly related species, and thus hybridization of relatively closely related species is advisable. Alternatively, preparation of an antibody preparation which exhibits less species specificity may be useful in expression cloning approaches.

#### III. Purified IL-170 protein

The predicted sequence of primate, e.g., human, IL-171 polypeptide sequence is shown in Table 1. Similarly, in Table 2, is provided primate, e.g., human, IL-175 sequence, and is assigned SEQ ID NO: 8. The peptide sequences allow preparation of peptides to generate antibodies to recognize such segments. Table 3 provides sequences of primate and murine IL-172; Table 4 provides sequence of primate and murine IL-173; Table 5 provides sequence of primate and murine IL-174; and Table 6 provides sequence of primate IL-176 and IL-177. Table 7 compares various IL-170 family members.

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As used herein, the terms "primate IL-170 protein" and "rodent IL-170 protein" shall encompass, when used in a protein context, a protein having amino acid sequences shown in Tables 1-5, or a significant fragment of such a protein. It also refers to a primate or rodent derived polypeptide which exhibits similar biological function or interacts with IL-170 protein specific binding components. These binding components, e.g., antibodies, typically bind to an IL-170 protein with high affinity, e.g., at least about 100 nM, usually better than about 30 nM, preferably better than about 10 nM, and more preferably at better than about 3 nM. Homologous proteins would be found in mammalian species other than rat or humans, e.g., mouse, primates, and in the herpes virus genome, e.g., ORF13. Non-mammalian species should also possess structurally or functionally related genes and proteins.

The term "polypeptide" as used herein includes a significant fragment or segment, and encompasses a stretch of amino acid residues of at least about 8 amino acids, generally at least 10 amino acids, more generally at least 12 amino acids, often at least 14 amino acids, more often at least 16 amino acids, typically at least 18 amino acids, more typically at least 20 amino acids, usually at least 22 amino acids, more usually at least 24 amino acids, preferably at least 26 amino acids, more preferably at least 28 amino acids, and, in particularly preferred embodiments, at least about 30 or more amino acids. The specific ends of such a segment will be at any combinations within the protein, preferably encompassing structural domains.

The term "binding composition" refers to molecules that bind with specificity to IL-170 protein, e.g., in a ligand-receptor type fashion, an antibody-antigen interaction, or compounds, e.g., proteins which specifically associate with IL-170 protein, e.g., in a natural physiologically relevant protein-protein interaction, either covalent or non-covalent. The molecule may be a polymer, or chemical reagent. No implication as to whether IL-170 protein is either the ligand

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or the receptor of a ligand-receptor interaction is represented, other than the interaction exhibit similar specificity, e.g., specific affinity. A functional analog may be a protein with structural modifications, or may be a wholly unrelated molecule, e.g., which has a molecular shape which interacts with the appropriate binding determinants. The proteins may serve as agonists or antagonists of a receptor, see, e.g., Goodman, et al. (eds. 1990) Goodman & Gilman's: The Pharmacological Bases of Therapeutics (8th ed.), Pergamon Press.

Solubility of a polypeptide or fragment depends upon the environment and the polypeptide. Many parameters affect polypeptide solubility, including temperature, electrolyte environment, size and molecular characteristics of the 15 polypeptide, and nature of the solvent. Typically, the temperature at which the polypeptide is used ranges from about 4° C to about 65° C. Usually the temperature at use is greater than about 18° C and more usually greater than about . 22° C. For diagnostic purposes, the temperature will usually 20 be about room temperature or warmer, but less than the denaturation temperature of components in the assay. For therapeutic purposes, the temperature will usually be body temperature, typically about 37° C for humans, though under certain situations the temperature may be raised or lowered in 25 situ or in vitro.

The electrolytes will usually approximate in situ physiological conditions, but may be modified to higher or lower ionic strength where advantageous. The actual ions may be modified, e.g., to conform to standard buffers used in physiological or analytical contexts.

The size and structure of the polypeptide should generally be in a substantially stable state, and usually not in a denatured state. The polypeptide may be associated with other polypeptides in a quaternary structure, e.g., to confer solubility, or associated with lipids or detergents in a manner which approximates natural lipid bilayer interactions.

The solvent will usually be a biologically compatible buffer, of a type used for preservation of biological activities, and will usually approximate a physiological solvent. Usually the solvent will have a neutral pH, typically between about 5 and 10, and preferably about 7.5. On some occasions, a detergent will be added, typically a mild non-denaturing one, e.g., CHS or CHAPS, or a low enough concentration as to avoid significant disruption of structural or physiological properties of the antigen.

10 Solubility is reflected by sedimentation measured in Svedberg units, which are a measure of the sedimentation velocity of a molecule under particular conditions. determination of the sedimentation velocity was classically performed in an analytical ultracentrifuge, but is typically 15 now performed in a standard ultracentrifuge. See, Freifelder (1982) Physical Biochemistry (2d ed.), W.H. Freeman; and Cantor and Schimmel (1980) Biophysical Chemistry, parts 1-3, W.H. Freeman & Co., San Francisco. As a crude determination, a sample containing a putatively soluble polypeptide is spun 20 in a standard full sized ultracentrifuge at about 50K rpm for about 10 minutes, and soluble molecules will remain in the supernatant. A soluble particle or polypeptide will typically be less than about 30S, more typically less than about 15S, usually less than about 10S, more usually less than about 6S, 25 and, in particular embodiments, preferably less than about 4S, and more preferably less than about 3S.

# IV. Making IL-170 protein; Mimetics

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DNA which encodes the IL-170 protein or fragments thereof can be obtained by chemical synthesis, screening cDNA libraries, or by screening genomic libraries prepared from a wide variety of cell lines or tissue samples.

This DNA can be expressed in a wide variety of host cells for the synthesis of a full-length protein or fragments which can in turn, for example, be used to generate polyclonal or monoclonal antibodies; for binding studies; for construction and expression of modified molecules; and for

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structure/function studies. Each antigen or its fragments can be expressed in host cells that are transformed or transfected with appropriate expression vectors. These molecules can be substantially purified to be free of protein or cellular contaminants, other than those derived from the recombinant host, and therefore are particularly useful in pharmaceutical compositions when combined with a pharmaceutically acceptable carrier and/or diluent. The antigen, or portions thereof, may be expressed as fusions with other proteins.

Expression vectors are typically self-replicating DNA or RNA constructs containing the desired antigen gene or its fragments, usually operably linked to suitable genetic control elements that are recognized in a suitable host cell. control elements are capable of effecting expression within a suitable host. The specific type of control elements necessary to effect expression will depend upon the eventual host cell used. Generally, the genetic control elements can include a prokaryotic promoter system or a eukaryotic promoter expression control system, and typically include a transcriptional promoter, an optional operator to control the onset of transcription, transcription enhancers to elevate the level of mRNA expression, a sequence that encodes a suitable ribosome binding site, and sequences that terminate transcription and translation. Expression vectors also usually contain an origin of replication that allows the vector to replicate independently of the host cell. Methods for amplifying vector copy number are also known, see, e.g., Kaufman, et al. (1985) Molec. and Cell. Biol. 5:1750-1759.

The vectors of this invention contain DNA which encodes an IL-170 protein, or a fragment thereof, typically encoding a biologically active polypeptide. The DNA can be under the control of a viral promoter and can encode a selection marker. This invention further contemplates use of such expression vectors which are capable of expressing eukaryotic cDNA coding for an IL-170 protein in a prokaryotic or eukaryotic host, where the vector is compatible with the host and where the eukaryotic cDNA coding for the antigen is inserted into the

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vector such that growth of the host containing the vector expresses the cDNA in question. Usually, expression vectors are designed for stable replication in their host cells or for amplification to greatly increase the total number of copies of the desirable gene per cell. It is not always necessary to require that an expression vector replicate in a host cell, e.g., it is possible to effect transient expression of the antigen or its fragments in various hosts using vectors that do not contain a replication origin that is recognized by the host cell. It is also possible to use vectors that cause integration of an IL-170 protein gene or its fragments into the host DNA by recombination, or to integrate a promoter which controls expression of an endogenous gene.

Vectors, as used herein, comprise plasmids, viruses, 15 bacteriophage, integratable DNA fragments, and other vehicles which enable the integration of DNA fragments into the genome of the host. Expression vectors are specialized vectors which contain genetic control elements that effect expression of operably linked genes. Plasmids are the most commonly used 20 form of vector but all other forms of vectors which serve an equivalent function and which are, or become, known in the art are suitable for use herein. See, e.g., Pouwels, et al. (1985 and Supplements) Cloning Vectors: A Laboratory Manual, Elsevier, N.Y., and Rodriquez, et al. (eds. 1988) Vectors: A 25 Survey of Molecular Cloning Vectors and Their Uses, Buttersworth, Boston, MA.

Transformed cells include cells, preferably mammalian, that have been transformed or transfected with vectors containing an IL-170 gene, typically constructed using recombinant DNA techniques. Transformed host cells usually express the antigen or its fragments, but for purposes of cloning, amplifying, and manipulating its DNA, do not need to express the protein. This invention further contemplates culturing transformed cells in a nutrient medium, thus permitting the protein to accumulate in the culture. The protein can be recovered, either from the culture or from the culture medium.

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For purposes of this invention, DNA sequences are operably linked when they are functionally related to each other. For example, DNA for a presequence or secretory leader is operably linked to a polypeptide if it is expressed as a preprotein or participates in directing the polypeptide to the cell membrane or in secretion of the polypeptide. A promoter is operably linked to a coding sequence if it controls the transcription of the polypeptide; a ribosome binding site is operably linked to a coding sequence if it is positioned to permit translation. Usually, operably linked means contiguous and in reading frame, however, certain genetic elements such as repressor genes are not contiguously linked but still bind to operator sequences that in turn control expression.

Suitable host cells include prokaryotes, lower eukaryotes, and higher eukaryotes. Prokaryotes include both gram negative and gram positive organisms, e.g., E. coli and B. subtilis. Lower eukaryotes include yeasts, e.g., S. cerevisiae and Pichia, and species of the genus Dictyostelium. Higher eukaryotes include established tissue culture cell lines from animal cells, both of non-mammalian origin, e.g., insect cells, and birds, and of mammalian origin, e.g., human, primates, and rodents.

Prokaryotic host-vector systems include a wide variety of vectors for many different species. As used herein, E. coli and its vectors will be used generically to include equivalent vectors used in other prokaryotes. A representative vector for amplifying DNA is pBR322 or many of its derivatives.

Vectors that can be used to express the IL-170 proteins or its fragments include, but are not limited to, such vectors as those containing the lac promoter (pUC-series); trp promoter (pBR322-trp); Ipp promoter (the pIN-series); lambda-pP or pR promoters (pOTS); or hybrid promoters such as ptac (pDR540).

See Brosius, et al. (1988) "Expression Vectors Employing Lambda-, trp-, lac-, and Ipp-derived Promoters", in Rodriguez and Denhardt (eds.) Vectors: A Survey of Molecular Cloning Vectors and Their Uses, Buttersworth, Boston, Chapter 10, pp. 205-236.

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Lower eukaryotes, e.g., yeasts and Dictyostelium, may be transformed with vectors encoding IL-170 proteins. purposes of this invention, the most common lower eukaryotic host is the baker's yeast, Saccharomyces cerevisiae. be used to generically represent lower eukaryotes although a number of other strains and species are also available. Yeast vectors typically consist of a replication origin (unless of the integrating type), a selection gene, a promoter, DNA encoding the desired protein or its fragments, and sequences for translation termination, polyadenylation, and transcription termination. Suitable expression vectors for yeast include such constitutive promoters as 3phosphoglycerate kinase and various other glycolytic enzyme gene promoters or such inducible promoters as the alcohol dehydrogenase 2 promoter or metallothionine promoter. Suitable vectors include derivatives of the following types: self-replicating low copy number (such as the YRp-series), self-replicating high copy number (such as the YEp-series); integrating types (such as the YIp-series), or minichromosomes (such as the YCp-series).

Higher eukaryotic tissue culture cells are the preferred host cells for expression of the functionally active IL-170 protein. In principle, many higher eukaryotic tissue culture cell lines are workable, e.g., insect baculovirus expression systems, whether from an invertebrate or vertebrate source. However, mammalian cells are preferred, in that the processing, both cotranslationally and posttranslationally. Transformation or transfection and propagation of such cells has become a routine procedure. Examples of useful cell lines include HeLa cells, Chinese hamster ovary (CHO) cell lines, baby rat kidney (BRK) cell lines, insect cell lines, bird cell lines, and monkey (COS) cell lines. Expression vectors for such cell lines usually include an origin of replication, a promoter, a translation initiation site, RNA splice sites (if genomic DNA is used), a polyadenylation site, and a transcription termination site. These vectors also usually contain a selection gene or amplification gene.

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expression vectors may be plasmids, viruses, or retroviruses carrying promoters derived, e.g., from such sources as from adenovirus, SV40, parvoviruses, vaccinia virus, or cytomegalovirus. Representative examples of suitable expression vectors include pCDNA1; pCD, see Okayama, et al. (1985) Mol. Cell Biol. 5:1136-1142; pMClneo Poly-A, see Thomas, et al. (1987) Cell 51:503-512; and a baculovirus vector such as pAC 373 or pAC 610, see O'Reilly, et al. (1992) Baculovirus Expression Vectors: A Laboratory Manual Freeman and Co., CRC Press, Boca Raton, Fla.

It will often be desired to express an IL-170 protein polypeptide in a system which provides a specific or defined glycosylation pattern. In this case, the usual pattern will be that provided naturally by the expression system. However, the pattern will be modifiable by exposing the polypeptide, e.g., an unglycosylated form, to appropriate glycosylating proteins introduced into a heterologous expression system. For example, the IL-170 protein gene may be co-transformed with one or more genes encoding mammalian or other glycosylating enzymes. Using this approach, certain mammalian glycosylation patterns will be achievable or approximated in prokaryote or other cells.

The IL-170 protein, or a fragment thereof, may be engineered to be phosphatidyl inositol (PI) linked to a cell membrane, but can be removed from membranes by treatment with a phosphatidyl inositol cleaving enzyme, e.g., phosphatidyl inositol phospholipase-C. This releases the antigen in a biologically active form, and allows purification by standard procedures of protein chemistry. See, e.g., Low (1989)

Biochim. Biophys. Acta 988:427-454; Tse, et al. (1985) Science 230:1003-1008; and Brunner, et al. (1991) J. Cell Biol. 114:1275-1283.

Now that the IL-170 protein has been characterized, fragments or derivatives thereof can be prepared by conventional processes for synthesizing peptides. These include processes such as are described in Stewart and Young (1984) Solid Phase Peptide Synthesis, Pierce Chemical Co.,

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Rockford, IL; Bodanszky and Bodanszky (1984) The Practice of Peptide Synthesis, Springer-Verlag, New York; and Bodanszky (1984) The Principles of Peptide Synthesis, Springer-Verlag, New York. For example, an azide process, an acid chloride process, an acid anhydride process, a mixed anhydride process, an active ester process (for example, p-nitrophenyl ester, N-hydroxysuccinimide ester, or cyanomethyl ester), a carbodiimidazole process, an oxidative-reductive process, or a dicyclohexylcarbodiimide (DCCD)/additive process can be used. Solid phase and solution phase syntheses are both applicable to the foregoing processes.

The IL-170 protein, fragments, or derivatives are suitably prepared in accordance with the above processes as typically employed in peptide synthesis, generally either by a so-called stepwise process which comprises condensing an amino acid to the terminal amino acid, one by one in sequence, or by coupling peptide fragments to the terminal amino acid. Amino groups that are not being used in the coupling reaction are typically protected to prevent coupling at an incorrect location.

If a solid phase synthesis is adopted, the C-terminal amino acid is bound to an insoluble carrier or support through its carboxyl group. The insoluble carrier is not particularly limited as long as it has a binding capability to a reactive carboxyl group. Examples of such insoluble carriers include halomethyl resins, such as chloromethyl resin or bromomethyl resin, hydroxymethyl resins, phenol resins, tertalkyloxycarbonyl-hydrazidated resins, and the like.

An amino group-protected amino acid is bound in sequence through condensation of its activated carboxyl group and the reactive amino group of the previously formed peptide or chain, to synthesize the peptide step by step. After synthesizing the complete sequence, the peptide is split off from the insoluble carrier to produce the peptide. This solid-phase approach is generally described by Merrifield, et al. (1963) in <u>J. Am. Chem. Soc.</u> 85:2149-2156.

The prepared protein and fragments thereof can be isolated and purified from the reaction mixture by means of peptide separation, for example, by extraction, precipitation, electrophoresis and various forms of chromatography, and the The IL-170 proteins of this invention can be obtained in varying degrees of purity depending upon its desired use. Purification can be accomplished by use of the protein purification techniques disclosed herein or by the use of the antibodies herein described in immunoabsorbant affinity 10 chromatography. This immunoabsorbant affinity chromatography is carried out by first linking the antibodies to a solid support and then contacting the linked antibodies with solubilized lysates of appropriate source cells, lysates of other cells expressing the protein, or lysates or supernatants of cells producing the IL-170 protein as a result of DNA 15 techniques, see below.

## V. Physical Variants

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This invention also encompasses proteins or peptides having substantial amino acid sequence homology with the amino acid sequence of the IL-170 protein. The variants include species or allelic variants.

Amino acid sequence homology, or sequence identity, is determined by optimizing residue matches, if necessary, by 25 introducing gaps as required. This changes when considering conservative substitutions as matches. Conservative substitutions typically include substitutions within the following groups: glycine, alanine; valine, isoleucine, leucine; aspartic acid, glutamic acid; asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, 30 tyrosine. Homologous amino acid sequences are typically intended to include natural allelic and interspecies variations in each respective protein sequence. homologous proteins or peptides will have from 25-100% homology (if gaps can be introduced), to 50-100% homology (if 35 conservative substitutions are included) with the amino acid sequence of the IL-170 protein. Homology measures will be at

least about 35%, generally at least 40%, more generally at least 45%, often at least 50%, more often at least 55%, typically at least 60%, more typically at least 65%, usually at least 70%, more usually at least 75%, preferably at least 80%, and more preferably at least 80%, and in particularly preferred embodiments, at least 85% or more. See also Needleham, et al. (1970) J. Mol. Biol. 48:443-453; Sankoff, et al. (1983) Chapter One in Time Warps, String Edits, and Macromolecules: The Theory and Practice of Sequence Comparison Addison-Wesley, Reading, MA; and software packages from IntelliGenetics, Mountain View, CA; and the University of Wisconsin Genetics Computer Group, Madison, WI.

The isolated DNA encoding an IL-170 protein can be readily modified by nucleotide substitutions, nucleotide deletions, nucleotide insertions, and inversions of nucleotide 15 These modifications result in novel DNA sequences stretches. which encode these antigens, their derivatives, or proteins having similar physiological, immunogenic, or antigenic These modified sequences can be used to produce activity. 20 mutant antigens or to enhance expression. Enhanced expression may involve gene amplification, increased transcription, increased translation, and other mechanisms. Such mutant IL-170 protein derivatives include predetermined or site-specific mutations of the respective protein or its fragments. 25 IL-170 protein encompasses a polypeptide otherwise falling within the homology definition of the murine IL-170 or human IL-170 protein as set forth above, but having an amino acid sequence which differs from that of IL-170 protein as found in nature, whether by way of deletion, substitution, or 30 insertion. In particular, "site specific mutant IL-170 protein generally includes proteins having significant homology with a protein having sequences of Tables 1-5, and as sharing various biological activities, e.g., antigenic or immunogenic, with those sequences, and in preferred 35 embodim nts contain most of the disclosed sequences. concepts apply to different IL-170 proteins, particularly

those found in various warm blooded animals, e.g., mammals and

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birds. As stated before, it is emphasized that descriptions are generally meant to encompass all IL-170 proteins, not limited to the mouse embodiment specifically discussed.

Although site specific mutation sites are predetermined, mutants need not be site specific. IL-170 protein mutagenesis can be conducted by making amino acid insertions or deletions. Substitutions, deletions, insertions, or any combinations may be generated to arrive at a final construct. Insertions include amino- or carboxy- terminal fusions. Random mutagenesis can be conducted at a target codon and the expressed mutants can then be screened for the desired activity. Methods for making substitution mutations at predetermined sites in DNA having a known sequence are well known in the art, e.g., by M13 primer mutagenesis or polymerase chain reaction (PCR) techniques. See also Sambrook, et al. (1989) and Ausubel, et al. (1987 and Supplements).

The mutations in the DNA normally should not place coding sequences out of reading frames and preferably will not create complementary regions that could hybridize to produce secondary mRNA structure such as loops or hairpins.

The present invention also provides recombinant proteins, e.g., heterologous fusion proteins using segments from these proteins. A heterologous fusion protein is a fusion of proteins or segments which are naturally not normally fused in the same manner. Thus, the fusion product of an immunoglobulin with an IL-170 polypeptide is a continuous protein molecule having sequences fused in a typical peptide linkage, typically made as a single translation product and exhibiting properties derived from each source peptide. A similar concept applies to heterologous nucleic acid sequences.

In addition, new constructs may be made from combining similar functional domains from other proteins. For example, antigen-binding or other segments may be "swapped" between different new fusion polypeptides or fragments. See, e.g., Cunningham, et al. (1989) <u>Science</u> 243:1330-1336; and O'Dowd,

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et al. (1988) <u>J. Biol. Chem.</u> 263:15985-15992. Thus, new chimeric polypeptides exhibiting new combinations of specificities will result from the functional linkage of biologically relevant domains and other functional domains.

The phosphoramidite method described by Beaucage and Carruthers (1981) <u>Tetra. Letts.</u> 22:1859-1862, will produce suitable synthetic DNA fragments. A double stranded fragment will often be obtained either by synthesizing the complementary strand and annealing the strand together under appropriate conditions or by adding the complementary strand using DNA polymerase with an appropriate primer sequence, e.g., PCR techniques.

## VI. Functional Variants

The blocking of physiological response to IL-170 proteins may result from the inhibition of binding of the antigen to its natural binding partner, e.g., through competitive inhibition. Thus, in vitro assays of the present invention will often use isolated protein, membranes from cells expressing a recombinant membrane associated IL-170 protein, soluble fragments comprising binding segments, or fragments attached to solid phase substrates. These assays will also allow for the diagnostic determination of the effects of either binding segment mutations and modifications, or protein mutations and modifications, e.g., analogs.

This invention also contemplates the use of competitive drug screening assays, e.g., where neutralizing antibodies to antigen or binding partner fragments compete with a test compound for binding to the protein. In this manner, the antibodies can be used to detect the presence of any polypeptide which shares one or more antigenic binding sites of the protein and can also be used to occupy binding sites on the protein that might otherwise interact with a binding partner.

35 Additionally, neutralizing antibodies against the IL-170 protein and soluble fragments of the antigen which contain a high affinity receptor binding site, can be used to inhibit

antigen function in tissues, e.g., tissues experiencing abnormal physiology.

"Derivatives" of the IL-170 antigens include amino acid sequence mutants, glycosylation variants, and covalent or aggregate conjugates with other chemical moieties. Covalent derivatives can be prepared by linkage of functionalities to groups which are found in the IL-170 amino acid side chains or at the N- or C- termini, by means which are well known in the art. These derivatives can include, without limitation, aliphatic esters or amides of the carboxyl terminus, or of 10 residues containing carboxyl side chains, 0-acyl derivatives of hydroxyl group-containing residues, and N-acyl derivatives of the amino terminal amino acid or amino-group containing residues, e.g., lysine or arginine. Acyl groups are selected from the group of alkyl-moieties including C3 to C18 normal 15 alkyl, thereby forming alkanoyl aroyl species. Covalent attachment to carrier proteins may be important when immunogenic moieties are haptens.

In particular, glycosylation alterations are included,
20 e.g., made by modifying the glycosylation patterns of a
polypeptide during its synthesis and processing, or in further
processing steps. Particularly preferred means for
accomplishing this are by exposing the polypeptide to
glycosylating enzymes derived from cells which normally
25 provide such processing, e.g., mammalian glycosylation
enzymes. Deglycosylation enzymes are also contemplated. Also
embraced are versions of the same primary amino acid sequence
which have other minor modifications, including phosphorylated
amino acid residues, e.g., phosphotyrosine, phosphoserine, or
30 phosphothreonine.

A major group of derivatives are covalent conjugates of the IL-170 protein or fragments thereof with other proteins or polypeptides. These derivatives can be synthesized in recombinant culture such as N- or C-terminal fusions or by the use of agents known in the art for their usefulness in crosslinking proteins through reactive side groups. Preferred antigen derivatization sites with cross-linking agents are at

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free amino groups, carbohydrate moieties, and cysteine residues.

Fusion polypeptides between the IL-170 proteins and other homologous or heterologous proteins are also provided. Homologous polypeptides may be fusions between different surface markers, resulting in, e.g., a hybrid protein exhibiting receptor binding specificity. Likewise, heterologous fusions may be constructed which would exhibit a combination of properties or activities of the derivative 10 proteins. Typical examples are fusions of a reporter polypeptide, e.g., luciferase, with a segment or domain of an antigen, e.g., a receptor-binding segment, so that the presence or location of the fused antigen may be easily determined. See, e.g., Dull, et al., U.S. Patent No. 15 4,859,609. Other gene fusion partners include bacterial ßgalactosidase, trpE, Protein A, ß-lactamase, alpha amylase, alcohol dehydrogenase, and yeast alpha mating factor.

The phosphoramidite method described by Beaucage and Carruthers (1981) <u>Tetra. Letts.</u> 22:1859-1862, will produce suitable synthetic DNA fragments. A double stranded fragment will often be obtained either by synthesizing the complementary strand and annealing the strand together under appropriate conditions or by adding the complementary strand using DNA polymerase with an appropriate primer sequence.

e.g., Godowski, et al. (1988) Science 241:812-816.

Such polypeptides may also have amino acid residues which have been chemically modified by phosphorylation, sulfonation, biotinylation, or the addition or removal of other moieties, particularly those which have molecular shapes similar to phosphate groups. In some embodiments, the modifications will be useful labeling reagents, or serve as purification targets, e.g., affinity ligands.

Fusion proteins will typically be made by either recombinant nucleic acid methods or by synthetic polypeptide methods. Techniques for nucleic acid manipulation and expression are described generally, for example, in Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual (2d)

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ed.), Vols. 1-3, Cold Spring Harbor Laboratory. Techniques for synthesis of polypeptides are described, for example, in Merrifield (1963) J. Amer. Chem. Soc. 85:2149-2156; Merrifield (1986) Science 232: 341-347; and Atherton, et al. (1989) Solid Phase Peptide Synthesis: A Practical Approach, IRL Press, Oxford.

This invention also contemplates the use of derivatives of the IL-170 proteins other than variations in amino acid sequence or glycosylation. Such derivatives may involve covalent or aggregative association with chemical moieties. These derivatives generally fall into the three classes: (1) salts, (2) side chain and terminal residue covalent modifications, and (3) adsorption complexes, for example with cell membranes. Such covalent or aggregative derivatives are useful as immunogens, as reagents in immunoassays, or in purification methods such as for affinity purification of antigens or other binding proteins. For example, an IL-170 antigen can be immobilized by covalent bonding to a solid support such as cyanogen bromide-activated Sepharose, by methods which are well known in the art, or adsorbed onto 20 polyolefin surfaces, with or without glutaraldehyde crosslinking, for use in the assay or purification of anti-IL-170 protein antibodies or its receptor or other binding partner. The IL-170 antigens can also be labeled with a detectable group, for example radioiodinated by the chloramine T procedure, covalently bound to rare earth chelates, or conjugated to another fluorescent moiety for use in diagnostic assays. Purification of IL-170 protein may be effected by immobilized antibodies or binding partners.

A solubilized IL-170 antigen or fragment of this invention can be used as an immunogen for the production of antisera or antibodies specific for the protein or fragments thereof. The purified antigen can be used to screen monoclonal antibodies or binding fragments prepared by 35 immunization with various forms of impure preparations containing the protein. In particular, the term "antibodies" also encompasses antigen binding fragments of natural

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antibodies. The purified IL-170 proteins can also be used as a reagent to detect any antibodies generated in response to the presence of elevated levels of the protein or cell fragments containing the antigen, both of which may be diagnostic of an abnormal or specific physiological or disease condition. Additionally, antigen fragments may also serve as immunogens to produce the antibodies of the present invention, as described immediately below. For example, this invention contemplates antibodies raised against amino acid sequences encoded by nucleotide sequences shown in Table 1 through 5, or fragments of proteins containing them. In particular, this invention contemplates antibodies having binding affinity to or being raised against specific fragments which are predicted to lie outside of the lipid bilayer.

The present invention contemplates the isolation of additional closely related species variants. Southern blot analysis established that similar genetic entities exist in other mammals, e.g., rat and human. It is likely that the IL-170 proteins are widespread in species variants, e.g., rodents, lagomorphs, carnivores, artiodactyla, perissodactyla, and primates.

The invention also provides means to isolate a group of related antigens displaying both distinctness and similarities in structure, expression, and function. Elucidation of many of the physiological effects of the antigens will be greatly accelerated by the isolation and characterization of distinct species variants. In particular, the present invention provides useful probes for identifying additional homologous genetic entities in different species.

The isolated genes will allow transformation of cells lacking expression of a corresponding IL-170 protein, e.g., either species types or cells which lack corresponding antigens and should exhibit negative background activity. Expression of transformed genes will allow isolation of antigenically pure cell lines, with defined or single specie variants. This approach will allow for more sensitive detection and discrimination of the physiological effects of

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IL-170 proteins. Subcellular fragments, e.g., cytoplasts or membrane fragments, can be isolated and used.

Dissection of the critical structural elements which effect the various physiological or differentiation functions provided by the proteins is possible using standard techniques of modern molecular biology, particularly in comparing members of the related class. See, e.g., the homolog-scanning mutagenesis technique described in Cunningham, et al. (1989) <a href="Science 243:1339-1336">Science 243:1339-1336</a>; and approaches used in O'Dowd, et al. (1988) <a href="J. Biol. Chem.">J. Biol. Chem.</a> 263:15985-15992; and Lechleiter, et al. (1990) <a href="EMBO J.">EMBO J.</a> 9:4381-4390.

In particular, functional domains or segments can be substituted between species variants to determine what structural features are important in both binding partner affinity and specificity, as well as signal transduction. An array of different variants will be used to screen for molecules exhibiting combined properties of interaction with different species variants of binding partners.

Antigen internalization may occur under certain circumstances, and interaction between intracellular 20 components and "extracellular" segments of proteins involved in interactions may occur. The specific segments of interaction of IL-170 protein with other intracellular components may be identified by mutagenesis or direct biochemical means, e.g., cross-linking or affinity methods. 25 Structural analysis by crystallographic or other physical methods will also be applicable. Further investigation of the mechanism of biological function will include study of associated components which may be isolatable by affinity methods or by genetic means, e.g., complementation analysis of 30 mutants.

Further study of the expression and control of IL-170 protein will be pursued. The controlling elements associated with the antigens may exhibit differential developmental, tissue specific, or other expression patterns. Upstream or downstream genetic regions, e.g., control elements, are of interest.

Structural studies of the antigen will lead to design of new variants, particularly analogs exhibiting agonist or antagonist properties on binding partners. This can be combined with previously described screening methods to isolate variants exhibiting desired spectra of activities.

Expression in other cell types will often result in glycosylation differences in a particular antigen. Various species variants may exhibit distinct functions based upon structural differences other than amino acid sequence. Differential modifications may be responsible for differential function, and elucidation of the effects are now made possible.

Thus, the present invention provides important reagents related to antigen-binding partner interaction. Although the foregoing description has focused primarily upon the murine IL-170 and human IL-170 protein, those of skill in the art will immediately recognize that the invention encompasses other antigens, e.g., mouse and other mammalian species or allelic variants, as well as variants thereof.

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### VII. Antibodies

Antibodies can be raised to the various IL-170 proteins, including species or allelic variants, and fragments thereof, both in their naturally occurring forms and in their recombinant forms. Additionally, antibodies can be raised to IL-170 proteins in either their active forms or in their inactive forms. Anti-idiotypic antibodies are also contemplated.

Antibodies, including binding fragments and single chain versions, against predetermined fragments of the antigens can be raised by immunization of animals with conjugates of the fragments with immunogenic proteins. Monoclonal antibodies are prepared from cells secreting the desired antibody. These antibodies can be screened for binding to normal or defective IL-170 proteins, or screened for agonistic or antagonistic activity, e.g., mediated through a binding partner. These monoclonal antibodies will usually bind with at least a  $K_{\rm D}$  of

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about 1 mM, more usually at least about 300  $\mu$ M, typically at least about 10  $\mu$ M, more typically at least about 30  $\mu$ M, preferably at least about 10  $\mu$ M, and more preferably at least about 3  $\mu$ M or better.

An IL-170 polypeptide that specifically binds to or that is specifically immunoreactive with an antibody, e.g., such as a polyclonal antibody, generated against a defined immunogen, e.g., such as an immunogen consisting of an amino acid sequence of SEQ ID NO: 3 or fragments thereof or a polypeptide generated from the nucleic acid of SEQ ID NO: 1 is typically determined in an immunoassay. Included within the metes and bounds of the present invention are those nucleic acid sequences described herein, including functional variants, that encode polypeptides that selectively bind to polyclonal antibodies generated against the prototypical IL-171 polypeptide as structurally and functionally defined herein. The immunoassay typically uses a polyclonal antiserum which was raised, e.g., to a protein of SEQ ID NO: 3 or 5. This antiserum is selected to have low crossreactivity against appropriate other IL-170 family members, preferably from the same species, and any such crossreactivity is removed by immunoabsorption prior to use in the immunoassay. Appropriate selective serum preparations can be isolated, and characterized.

In order to produce antisera for use in an immunoassay, the protein, e.g., of SEQ ID NO: 5, is isolated as described herein. For example, recombinant protein may be produced in a mammalian cell line. An appropriate host, e.g., an inbred strain of mice such as Balb/c, is immunized with the protein of SEQ ID NO: 5 using a standard adjuvant, such as Freund's adjuvant, and a standard mouse immunization protocol (see Harlow and Lane). Alternatively, a substantially full length synthetic peptide derived from the sequences disclosed herein can be used as an immunogen. Polyclonal sera are collected and titered against the immunogen protein in an immunoassay, e.g., a solid phase immunoassay with the immunogen immobilized on a solid support. Polyclonal antisera with a titer of 104

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or greater are selected and tested for their cross reactivity against other IL-170 family members, e.g., IL-171, IL-172, IL-173, IL-174, IL-175, IL-176, or IL-177, using a competitive binding immunoassay such as the one described in Harlow and Lane, supra, at pages 570-573. Preferably at least two IL-170 family members are used in this determination in conjunction with the target. These IL-170 family members can be produced as recombinant proteins and isolated using standard molecular biology and protein chemistry techniques as described herein. Thus, antibody preparations can be identified or produced having desired selectivity or specificity for subsets of IL-170 family members.

Immunoassays in the competitive binding format can be used for the crossreactivity determinations. For example, the protein of SEQ ID NO: 5 can be immobilized to a solid support. Proteins added to the assay compete with the binding of the antisera to the immobilized antigen. The ability of the above proteins to compete with the binding of the antisera to the immobilized protein is compared to the protein of SEQ ID NO:

5. The percent crossreactivity for the above proteins is calculated, using standard calculations. Those antisera with less than 10% crossreactivity with each of the proteins listed above are selected and pooled. The cross-reacting antibodies are then removed from the pooled antisera by immunoabsorption with the above-listed proteins.

The immunoabsorbed and pooled antisera are then used in a competitive binding immunoassay as described above to compare a second protein to the immunogen protein. In order to make this comparison, the two proteins are each assayed at a wide range of concentrations and the amount of each protein required to inhibit 50% of the binding of the antisera to the immobilized protein is determined. If the amount of the second protein required is less than twice the amount of the protein of, e.g., SEQ ID NO: 5 that is required, then the second protein is said to specifically bind to an antibody generated to the immunogen.

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The antibodies, including antigen binding fragments, of this invention can have significant diagnostic or therapeutic value. They can be potent antagonists that bind to a binding partner and inhibit antigen binding or inhibit the ability of an antigen to elicit a biological response. They also can be useful as non-neutralizing antibodies and can be coupled to toxins or radionuclides so that when the antibody binds to the antigen, a cell expressing it, e.g., on its surface, is killed. Further, these antibodies can be conjugated to drugs or other therapeutic agents, either directly or indirectly by means of a linker, and may effect drug targeting.

The antibodies of this invention can also be useful in diagnostic applications. As capture or non-neutralizing antibodies, they can be screened for ability to bind to the antigens without inhibiting binding by a partner. As neutralizing antibodies, they can be useful in competitive binding assays. They will also be useful in detecting or quantifying IL-170 protein or its binding partners. See, e.g., Chan (ed. 1987) Immunoassay: A Practical Guide Academic Press, Orlando, Fla.; Ngo (ed. 1988) Nonisotopic Immunoassay Plenum Press, NY; and Price and Newman (eds. 1991) Principles and Practice of Immunoassay Stockton Press, NY.

Antigen fragments may be joined to other materials, particularly polypeptides, as fused or covalently joined polypeptides to be used as immunogens. An antigen and its fragments may be fused or covalently linked to a variety of immunogens, such as keyhole limpet hemocyanin, bovine serum albumin, tetanus toxoid, etc. See Microbiology, Hoeber Medical Division, Harper and Row, 1969; Landsteiner (1962) Specificity of Serological Reactions, Dover Publications, New York, and Williams, et al. (1967) Methods in Immunology and Immunochemistry, Vol. 1, Academic Press, New York, for descriptions of methods of preparing polyclonal antisera. A typical method involves hyperimmunization of an animal with an antigen. The blood of the animal is then collected shortly after the repeated immunizations and the gamma globulin is isolated.

In some instances, it is desirable to prepare monoclonal antibodies from various mammalian hosts, such as mice, rodents, primates, humans, etc. Description of techniques for preparing such monoclonal antibodies may be found in, e.g., Stites, et al. (eds.) Basic and Clinical Immunology (4th ed.), 5 Lange Medical Publications, Los Altos, CA, and references cited therein; Harlow and Lane (1988) Antibodies: A Laboratory Manual, CSH Press; Goding (1986) Monoclonal Antibodies: Principles and Practice (2d ed.) Academic Press, New York; and particularly in Kohler and Milstein (1975) in 10 Nature 256: 495-497, which discusses one method of generating monoclonal antibodies. Summarized briefly, this method involves injecting an animal with an immunogen. The animal is then sacrificed and cells taken from its spleen, which are then fused with myeloma cells. The result is a hybrid cell or 15 "hybridoma" that is capable of reproducing in vitro. population of hybridomas is then screened to isolate individual clones, each of which secrete a single antibody species to the immunogen. In this manner, the individual antibody species obtained are the products of immortalized and 20 cloned single B cells from the immune animal generated in response to a specific site recognized on the immunogenic substance.

Other suitable techniques involve in vitro exposure of lymphocytes to the antigenic polypeptides or alternatively to 25 selection of libraries of antibodies in phage or similar vectors. See, Huse, et al. (1989) "Generation of a Large Combinatorial Library of the Immunoglobulin Repertoire in Phage Lambda, \* Science 246:1275-1281; and Ward, et al. (1989) Nature 341:544-546. The polypeptides and antibodies of the 30 present invention may be used with or without modification, including chimeric or humanized antibodies. Frequently, the polypeptides and antibodies will be labeled by joining, either covalently or non-covalently, a substance which provides for a detectable signal. A wide variety of labels and conjugation 35 techniques are known and are reported extensively in both the scientific and patent literature. Suitable labels include

radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent moieties, chemiluminescent moieties, magnetic particles, and the like. Patents, teaching the use of such labels include U.S. Patent Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241. Also, recombinant immunoglobulins may be produced, see Cabilly, U.S. Patent No. 4,816,567.

The antibodies of this invention can also be used for affinity chromatography in isolating the protein. Columns can be prepared where the antibodies are linked to a solid support, e.g., particles, such as agarose, Sephadex, or the like, where a cell lysate may be passed through the column, the column washed, followed by increasing concentrations of a mild denaturant, whereby the purified IL-170 protein will be released.

The antibodies may also be used to screen expression libraries for particular expression products. Usually the antibodies used in such a procedure will be labeled with a moiety allowing easy detection of presence of antigen by antibody binding.

Antibodies raised against each IL-170 protein will also be useful to raise anti-idiotypic antibodies. These will be useful in detecting or diagnosing various immunological conditions related to expression of the respective antigens.

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# VIII. Uses

The present invention provides reagents which will find use in diagnostic applications as described elsewhere herein, e.g., in the general description for physiological or developmental abnormalities, or below in the description of kits for diagnosis.

This invention also provides reagents with significant therapeutic value. The IL-170 protein (naturally occurring or recombinant), fragments thereof, and antibodies thereto, along with compounds identified as having binding affinity to IL-170 protein, should be useful in the treatment of conditions associated with abnormal physiology or development, including

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abnormal proliferation, e.g., cancerous conditions, or degenerative conditions. Abnormal proliferation, regeneration, degeneration, and atrophy may be modulated by appropriate therapeutic treatment using the compositions provided herein. For example, a disease or disorder associated with abnormal expression or abnormal signaling by an IL-170 antigen should be a likely target for an agonist or antagonist of the protein.

Other abnormal developmental conditions are known in the cell types shown to possess IL-170 antigen mRNA by Northern blot analysis. See Berkow (ed.) The Merck Manual of Diagnosis and Therapy, Merck & Co., Rahway, N.J.; and Thorn, et al. Harrison's Principles of Internal Medicine, McGraw-Hill, N.Y. These problems may be susceptible to prevention or treatment using compositions provided herein.

Recombinant antibodies which bind to IL-170 can be purified and then administered to a patient. These reagents can be combined for therapeutic use with additional active or inert ingredients, e.g., in conventional pharmaceutically acceptable carriers or diluents, e.g., immunogenic adjuvants, along with physiologically innocuous stabilizers and excipients. These combinations can be sterile filtered and placed into dosage forms as by lyophilization in dosage vials or storage in stabilized aqueous preparations. This invention also contemplates use of antibodies or binding fragments thereof, including forms which are not complement binding.

Screening using IL-170 for binding partners or compounds having binding affinity to IL-170 antigen can be performed, including isolation of associated components. Subsequent biological assays can then be utilized to determine if the compound has intrinsic biological activity and is therefore an agonist or antagonist in that it blocks an activity of the antigen. This invention further contemplates the therapeutic use of antibodies to IL-170 protein as antagonists. This approach should be particularly useful with other IL-170 protein species variants.

The quantities of reagents necessary for effective therapy will depend upon many different factors, including means of administration, target site, physiological state of the patient, and other medicants administered. treatment dosages should be titrated to optimize safety and efficacy. Typically, dosages used in vitro may provide useful guidance in the amounts useful for in situ administration of these reagents. Animal testing of effective doses for treatment of particular disorders will provide further 10 predictive indication of human dosage. Various considerations are described, e.g., in Gilman, et al. (eds. 1990) Goodman and Gilman's: The Pharmacological Bases of Therapeutics, 8th Ed., Pergamon Press; and Remington's Pharmaceutical Sciences, 17th ed. (1990), Mack Publishing Co., Easton, Penn. Methods for 15 administration are discussed therein and below, e.g., for oral, intravenous, intraperitoneal, or intramuscular administration, transdermal diffusion, and others. See also Langer (1990) Science 249:1527-1533. Pharmaceutically acceptable carriers will include water, saline, buffers, and 20 other compounds described, e.g., in the Merck Index, Merck & Co., Rahway, New Jersey. Dosage ranges would ordinarily be expected to be in amounts lower than 1 mM concentrations, typically less than about 10  $\mu$ M concentrations, usually less than about 100 nM, preferably less than about 10 pM 25 (picomolar), and most preferably less than about 1 fM (femtomolar), with an appropriate carrier. Slow release formulations, or a slow release apparatus will often be utilized for continuous administration.

or its fragments, antagonists, and agonists, may be administered directly to the host to be treated or, depending on the size of the compounds, it may be desirable to conjugate them to carrier proteins such as ovalbumin or serum albumin prior to their administration. Therapeutic formulations may be administered in any conventional dosage formulation. While it is possible for the active ingredient to be administered alone, it is preferable to present it as a pharmaceutical

formulation. Formulations typically comprise at least one active ingredient, as defined above, together with one or more acceptable carriers thereof. Each carrier should be both pharmaceutically and physiologically acceptable in the sense of being compatible with the other ingredients and not injurious to the patient. Formulations include those suitable for oral, rectal, nasal, or parenteral (including subcutaneous, intramuscular, intravenous and intradermal) administration. The formulations may conveniently be 10 presented in unit dosage form and may be prepared by any methods well known in the art of pharmacy. See, e.g., Gilman, et al. (eds. 1990) Goodman and Gilman's: The Pharmacological Bases of Therapeutics, 8th Ed., Pergamon Press, Parrytown, NY; Remington's Pharmaceutical Sciences, 17th ed. (1990) Mack Publishing Co., Easton, Penn.; Avis, et al. (eds. 1993) 15 Pharmaceutical Dosage Forms: Parenteral Medications 2d ed., Dekker, NY; Lieberman, et al. (eds. 1990) Pharmaceutical Dosage Forms: Tablets 2d ed., Dekker, NY; and Lieberman, et al. (eds. 1990) Pharmaceutical Dosage Forms: Disperse Systems 20 Dekker, NY. The therapy of this invention may be combined with or used in association with other therapeutic, including cytokine, reagents.

Both the naturally occurring and the recombinant forms of the IL-170 proteins of this invention are particularly useful 25 in kits and assay methods which are capable of screening compounds for binding activity to the proteins. Several methods of automating assays have been developed in recent years so as to permit screening of tens of thousands of compounds in a short period. See, e.g., Fodor, et al. (1991) 30 Science 251:767-773, which describes means for testing of binding affinity by a plurality of defined polymers synthesized on a solid substrate. The development of suitable assays can be greatly facilitated by the availability of large amounts of purified, soluble IL-170 protein as provided by this invention. 35

This invention is particularly useful for screening compounds by using recombinant antigen in any of a variety of

drug screening techniques. The advantages of using a recombinant protein in screening for specific ligands include: (a) improved renewable source of the antigen from a specific source; (b) potentially greater number of antigen molecules per cell giving better signal to noise ratio in assays; and (c) species variant specificity (theoretically giving greater biological and disease specificity). The purified protein may be tested in numerous assays, typically in vitro assays, which evaluate biologically relevant responses. See, e.g., Coligan Current Protocols in Immunology; Hood, et al. Immunology Benjamin/Cummings; Paul (ed.) Fundamental Immunology; and Methods in Enzymology Academic Press.

One method of drug screening utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant DNA molecules expressing the IL-170 antigens. 15 Cells may be isolated which express an antigen in isolation from other functionally equivalent antigens. Such cells, either in viable or fixed form, can be used for standard protein-protein binding assays. See also, Parce, et al. (1989) Science 246:243-247; and Owicki, et al. (1990) Proc. 20 Nat'l Acad. Sci. USA 87:4007-4011, which describe sensitive methods to detect cellular responses. Competitive assays are particularly useful, where the cells (source of IL-170 protein) are contacted and incubated with a labeled binding partner or antibody having known binding affinity to the 25 ligand, such as  $^{125}I$ -antibody, and a test sample whose binding affinity to the binding composition is being measured. bound and free labeled binding compositions are then separated to assess the degree of antigen binding. The amount of test compound bound is inversely proportional to the amount of 30 labeled receptor binding to the known source. Any one of numerous techniques can be used to separate bound from free antigen to assess the degree of binding. This separation step could typically involve a procedure such as adhesion to filters followed by washing, adhesion to plastic followed by washing, or centrifugation of the cell membranes. Viable cells could also be used to screen for the effects of drugs on

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IL-170 protein mediated functions, e.g., second messenger levels, i.e., Ca<sup>++</sup>; cell proliferation; inositol phosphate pool changes; and others. Some detection methods allow for elimination of a separation step, e.g., a proximity sensitive detection system. Calcium sensitive dyes will be useful for detecting Ca<sup>++</sup> levels, with a fluorimeter or a fluorescence cell sorting apparatus.

Another method utilizes membranes from transformed eukaryotic or prokaryotic host cells as the source of the IL-170 protein. These cells are stably transformed with DNA vectors directing the expression of a membrane associated IL-170 protein, e.g., an engineered membrane bound form. Essentially, the membranes would be prepared from the cells and used in any receptor/ligand type binding assay such as the competitive assay set forth above.

Still another approach is to use solubilized, unpurified or solubilized, purified IL-170 protein from transformed eukaryotic or prokaryotic host cells. This allows for a "molecular" binding assay with the advantages of increased specificity, the ability to automate, and high drug test throughput.

Another technique for drug screening involves an approach which provides high throughput screening for compounds having suitable binding affinity to IL-170 and is described in detail in Geysen, European Patent Application 84/03564, published on September 13, 1984. First, large numbers of different small peptide test compounds are synthesized on a solid substrate, e.g., plastic pins or some other appropriate surface, see Fodor, et al. (1991). Then all the pins are reacted with solubilized, unpurified or solubilized, purified IL-170 binding composition, and washed. The next step involves detecting bound binding composition.

Rational drug design may also be based upon structural studies of the molecular shapes of the IL-170 protein and other effectors or analogs. Effectors may be other proteins which mediate other functions in response to antigen binding, or other proteins which normally interact with the antigen.

One means for determining which sites interact with specific other proteins is a physical structure determination, e.g., xray crystallography or 2 dimensional NMR techniques. These will provide guidance as to which amino acid residues form molecular contact regions. For a detailed description of protein structural determination, see, e.g., Blundell and Johnson (1976) Protein Crystallography, Academic Press, New York.

Purified IL-170 protein can be coated directly onto 10 plates for use in the aforementioned drug screening techniques. However, non-neutralizing antibodies to these ligands can be used as capture antibodies to immobilize the respective ligand on the solid phase.

#### 15 IX. Kits

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This invention also contemplates use of IL-170 proteins, fragments thereof, peptides, and their fusion products in a variety of diagnostic kits and methods for detecting the presence of a binding composition. Typically the kit will 20 have a compartment containing either a defined IL-170 peptide or gene segment or a reagent which recognizes one or the other, e.g., antigen fragments or antibodies.

A kit for determining the binding affinity of a test compound to an IL-170 protein would typically comprise a test compound; a labeled compound, for example an antibody having known binding affinity for the antigen; a source of IL-170 protein (naturally occurring or recombinant); and a means for separating bound from free labeled compound, such as a solid phase for immobilizing the antigen. Once compounds are 30 screened, those having suitable binding affinity to the antigen can be evaluated in suitable biological assays, as are well known in the art, to determine whether they exhibit similar biological activities to the natural antigen. availability of recombinant IL-170 protein polypeptides also provide well defined standards for calibrating such assays.

A preferred kit for determining the concentration of, for example, an IL-170 protein in a sample would typically

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comprise a labeled compound, e.g., antibody, having known binding affinity for the antigen, a source of antigen (naturally occurring or recombinant) and a means for separating the bound from free labeled compound, for example, a solid phase for immobilizing the IL-170 protein. Compartments containing reagents, and instructions, will normally be provided.

One method for determining the concentration of IL-170 protein in a sample would typically comprise the steps of: (1) preparing membranes from a sample comprised of a membrane bound IL-170 protein source; (2) washing the membranes and suspending them in a buffer; (3) solubilizing the antigen by incubating the membranes in a culture medium to which a suitable detergent has been added; (4) adjusting the detergent concentration of the solubilized antigen; (5) contacting and incubating said dilution with radiolabeled antibody to form complexes; (6) recovering the complexes such as by filtration through polyethyleneimine treated filters; and (7) measuring the radioactivity of the recovered complexes.

Antibodies, including antigen binding fragments, specific for the IL-170 protein or fragments are useful in diagnostic applications to detect the presence of elevated levels of IL-170 protein and/or its fragments. Such diagnostic assays can employ lysates, live cells, fixed cells, immunofluorescence, cell cultures, body fluids, and further can involve the detection of antigens related to the protein in serum, or the like. Diagnostic assays may be homogeneous (without a separation step between free reagent and protein-protein complex) or heterogeneous (with a separation step). Various commercial assays exist, such as radioimmunoassay (RIA), enzyme-linked immunosorbent assay (ELISA), enzyme immunoassay (EIA), enzyme-multiplied immunoassay technique (EMIT), substrate-labeled fluorescent immunoassay (SLFIA), and the like. For example, unlabeled antibodies can be employed by using a second antibody which is labeled and which recognizes the antibody to an IL-170 protein or to a particular fragment thereof. Similar assays have also been extensively discussed

in the literature. See, e.g., Harlow and Lane (1988) Antibodies: A Laboratory Manual, CSH.

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Anti-idiotypic antibodies may have similar use to diagnose presence of antibodies against an IL-170 protein, as such may be diagnostic of various abnormal states. example, overproduction of IL-170 protein may result in production of various immunological reactions which may be diagnostic of abnormal physiological states, particularly in proliferative cell conditions such as cancer or abnormal differentiation.

Frequently, the reagents for diagnostic assays are supplied in kits, so as to optimize the sensitivity of the assay. For the subject invention, depending upon the nature of the assay, the protocol, and the label, either labeled or unlabeled antibody, or labeled IL-170 protein is provided. This is usually in conjunction with other additives, such as buffers, stabilizers, materials necessary for signal production such as substrates for enzymes, and the like. Preferably, the kit will also contain instructions for proper use and disposal of the contents after use. Typically the kit has compartments for each useful reagent. Desirably, the reagents are provided as a dry lyophilized powder, where the reagents may be reconstituted in an aqueous medium providing appropriate concentrations of reagents for performing the assay.

Any of the aforementioned constituents of the drug screening and the diagnostic assays may be used without modification or may be modified in a variety of ways. For example, labeling may be achieved by covalently or noncovalently joining a moiety which directly or indirectly provides a detectable signal. In any of these assays, the antigen, test compound, IL-170 protein, or antibodies thereto can be labeled either directly or indirectly. Possibilities for direct labeling include label groups: radiolabels such as  $125_{\text{I}}$ , enzymes (U.S. Pat. No. 3,645,090) such as peroxidase and alkaline phosphatase, and fluorescent labels (U.S. Pat. No. 3,940,475) capable of monitoring the change in fluorescence

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intensity, wavelength shift, or fluorescence polarization. Possibilities for indirect labeling include biotinylation of one constituent followed by binding to avidin coupled to one of the above label groups.

There are also numerous methods of separating the bound from the free antigen, or alternatively the bound from the free test compound. The IL-170 protein can be immobilized on various matrixes followed by washing. Suitable matrixes include plastic such as an ELISA plate, filters, and beads. 10 Methods of immobilizing the IL-170 protein to a matrix include, without limitation, direct adhesion to plastic, use of a capture antibody, chemical coupling, and biotin-avidin. The last step in this approach involves the precipitation of protein-protein complex by any of several methods including those utilizing, e.g., an organic solvent such as polyethylene glycol or a salt such as ammonium sulfate. Other suitable separation techniques include, without limitation, the fluorescein antibody magnetizable particle method described in Rattle, et al. (1984) Clin. Chem. 30:1457-1461, and the double antibody magnetic particle separation as described in U.S. Pat. No. 4,659,678.

The methods for linking proteins or their fragments to the various labels have been extensively reported in the literature and do not require detailed discussion here. Many of the techniques involve the use of activated carboxyl groups either through the use of carbodiimide or active esters to form peptide bonds, the formation of thioethers by reaction of a mercapto group with an activated halogen such as chloroacetyl, or an activated olefin such as maleimide, for linkage, or the like. Fusion proteins will also find use in these applications.

Another diagnostic aspect of this invention involves use of oligonucleotide or polynucleotide sequences taken from the sequence of an IL-170 protein. These sequences can be used as probes for detecting levels of antigen message in samples from patients suspected of having an abnormal condition, e.g., cancer or developmental problem. The preparation of both RNA

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and titered against the immunogen protein in an immunoassay, e.g., a solid phase immunoassay with the immunogen immobilized on a solid support. Polyclonal antisera with a titer of 10<sup>4</sup> or greater are selected and tested for their cross reactivity against other IL-170 family members, e.g., IL-171, IL-172, IL-173, IL-174, IL-175, IL-176, or IL-177, using a competitive binding immunoassay such as the one described in Harlow and Lane, supra, at pages 570-573. Preferably at least two IL-170 family members are used in this determination in conjunction with the target. These IL-170 family members can be produced as recombinant proteins and isolated using standard molecular biology and protein chemistry techniques as described herein. Thus, antibody preparations can be identified or produced having desired selectivity or specificity for subsets of IL-170 family members.

Immunoassays in the competitive binding format can be used for the crossreactivity determinations. For example, the protein of SEQ ID NO: 5 can be immobilized to a solid support. Proteins added to the assay compete with the binding of the antisera to the immobilized antigen. The ability of the above proteins to compete with the binding of the antisera to the immobilized protein is compared to the protein of SEQ ID NO: 5. The percent crossreactivity for the above proteins is calculated, using standard calculations. Those antisera with less than 10% crossreactivity with each of the proteins listed above are selected and pooled. The cross-reacting antibodies are then removed from the pooled antisera by immunoabsorption with the above-listed proteins.

The immunoabsorbed and pooled antisera are then used in a competitive binding immunoassay as described above to compare a second protein to the immunogen protein. In order to make this comparison, the two proteins are each assayed at a wide range of concentrations and the amount of each protein required to inhibit 50% of the binding of the antisera to the immobilized protein is determined. If the amount of the second protein required is less than twice the amount of the protein of, e.g., SEQ ID NO: 5 that is required, then the

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second protein is said to specifically bind to an antibody generated to the immunogen.

The antibodies, including antigen binding fragments, of this invention can have significant diagnostic or therapeutic value. They can be potent antagonists that bind to a binding partner and inhibit antigen binding or inhibit the ability of an antigen to elicit a biological response. They also can be useful as non-neutralizing antibodies and can be coupled to toxins or radionuclides so that when the antibody binds to the antigen, a cell expressing it, e.g., on its surface, is killed. Further, these antibodies can be conjugated to drugs or other therapeutic agents, either directly or indirectly by means of a linker, and may effect drug targeting.

The antibodies of this invention can also be useful in diagnostic applications. As capture or non-neutralizing antibodies, they can be screened for ability to bind to the antigens without inhibiting binding by a partner. As neutralizing antibodies, they can be useful in competitive binding assays. They will also be useful in detecting or quantifying IL-170 protein or its binding partners. See, e.g., Chan (ed. 1987) Immunoassay: A Practical Guide Academic Press, Orlando, Fla.; Ngo (ed. 1988) Nonisotopic Immunoassay Plenum Press, NY; and Price and Newman (eds. 1991) Principles and Practice of Immunoassay Stockton Press, NY.

Antigen fragments may be joined to other materials, particularly polypeptides, as fused or covalently joined polypeptides to be used as immunogens. An antigen and its fragments may be fused or covalently linked to a variety of immunogens, such as keyhole limpet hemocyanin, bovine serum albumin, tetanus toxoid, etc. See Microbiology, Hoeber Medical Division, Harper and Row, 1969; Landsteiner (1962) Specificity of Serological Reactions, Dover Publications, New York, and Williams, et al. (1967) Methods in Immunology and Immunochemistry, Vol. 1, Academic Press, New York, for descriptions of methods of preparing polyclonal antisera. A typical method involves hyperimmunization of an animal with an antigen. The blood of the animal is then collected shortly

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after the repeated immunizations and the gamma globulin is isolated.

In some instances, it is desirable to prepare monoclonal antibodies from various mammalian hosts, such as mice, rodents, primates, humans, etc. Description of techniques for preparing such monoclonal antibodies may be found in, e.g., Stites, et al. (eds.) Basic and Clinical Immunology (4th ed.), Lange Medical Publications, Los Altos, CA, and references cited therein; Harlow and Lane (1988) Antibodies: A Laboratory Manual, CSH Press; Goding (1986) Monoclonal Antibodies: Principles and Practice (2d ed.) Academic Press, New York; and particularly in Kohler and Milstein (1975) in Nature 256: 495-497, which discusses one method of generating monoclonal antibodies. Summarized briefly, this method involves injecting an animal with an immunogen. The animal is 15 then sacrificed and cells taken from its spleen, which are then fused with myeloma cells. The result is a hybrid cell or "hybridoma" that is capable of reproducing in vitro. population of hybridomas is then screened to isolate individual clones, each of which secrete a single antibody 20 species to the immunogen. In this manner, the individual antibody species obtained are the products of immortalized and cloned single B cells from the immune animal generated in response to a specific site recognized on the immunogenic 25 substance.

Other suitable techniques involve in vitro exposure of lymphocytes to the antigenic polypeptides or alternatively to selection of libraries of antibodies in phage or similar vectors. See, Huse, et al. (1989) "Generation of a Large Combinatorial Library of the Immunoglobulin Repertoire in Phage Lambda," Science 246:1275-1281; and Ward, et al. (1989) Nature 341:544-546. The polypeptides and antibodies of the present invention may be used with or without modification, including chimeric or humanized antibodies. Frequently, the polypeptides and antibodies will be labeled by joining, either covalently or non-covalently, a substance which provides for a detectable signal. A wide variety of labels and conjugation

techniques are known and are reported extensively in both the scientific and patent literature. Suitable labels include radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent moieties, chemiluminescent moieties, magnetic particles, and the like. Patents, teaching the use of such labels include U.S. Patent Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241. Also, recombinant immunoglobulins may be produced, see Cabilly, U.S. Patent No. 4,816,567.

The antibodies of this invention can also be used for affinity chromatography in isolating the protein. Columns can be prepared where the antibodies are linked to a solid support, e.g., particles, such as agarose, Sephadex, or the like, where a cell lysate may be passed through the column, the column washed, followed by increasing concentrations of a mild denaturant, whereby the purified IL-170 protein will be released.

The antibodies may also be used to screen expression libraries for particular expression products. Usually the antibodies used in such a procedure will be labeled with a moiety allowing easy detection of presence of antigen by antibody binding.

Antibodies raised against each IL-170 protein will also be useful to raise anti-idiotypic antibodies. These will be useful in detecting or diagnosing various immunological conditions related to expression of the respective antigens.

# VIII. Uses

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The present invention provides reagents which will find use in diagnostic applications as described elsewhere herein, e.g., in the general description for physiological or developmental abnormalities, or below in the description of kits for diagnosis.

This invention also provides reagents with significant therapeutic value. The IL-170 protein (naturally occurring or recombinant), fragments thereof, and antibodies thereto, along with compounds identified as having binding affinity to IL-170

III. Biochemical Characterization of IL-170 proteins An IL-170 protein is expressed in heterologous cells, e.g., the native form or a recombinant form displaying the FLAG peptide at the carboxy terminus. See, e.g., Crowe, et al. (1992) OIAexpress: The High Level Expression and Protein Purification System QIAGEN, Inc. Chatsworth, CA; and Hopp, et al. (1988) Bio/Technology 6:1204-1210. These two forms are introduced into expression vectors, e.g., pME18S or pEE12, and subsequently transfected into appropriate cells, e.g., COS-7 or NSO cells, respectively. Electroporated cells are 10 cultivated, e.g., for 48 hours in RPMI medium supplemented with 10% Fetal Calf Serum. Cells are then incubated with 35s-Met and <sup>35</sup>S-Cys in order to label cellular proteins. Comparison of the proteins under reducing conditions on SDS-PAGE should show that cells transfected with full length 15 clones should secret a polypeptide of the appropriate size, e.g., about 15,000 daltons. Treatment with endoglycosidases will demonstrate whether there are N-glycosylated forms.

In order to produce larger quantities of native proteins, stable transformants of NSO cells can be prepared according to the methodology developed by Celltech (Slough, Berkshire, UK; International Patent Applications WO86/05807, WO87/04462, WO89/01036, and WO89/10404).

Typically, 1 liter of supernatant containing human IL-171 or IL-171-FLAG is passed, e.g., on a 60 ml column of Zn++ ions grafted to a Chelating Sepharose Fast Flow matrix (Pharmacia, Upsalla, Sweden). After washing with 10 volumes of binding buffer (His-Bind Buffer kit, Novagen, Madison, WI), the

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proteins retained by the metal ions are eluted with a gradient of 20-100 mM Imidazole. The content of human IL-170-FLAG in the eluted fractions is determined by dot blot using the anti-FLAG monoclonal antibody M2 (Eastman Kodak, New Haven, CT), whereas the content of human IL-172 is assessed, e.g., by silver staining of non-reducing SDS-PAGE. The IL-170 containing fractions are then pooled and dialyzed against PBS, and are either used in biological assays or further purified, e.g., by anion exchange HPLC on a DEAE column. A third step of gel filtration chromatography may be performed on a SUPERDEX G-75 HRD30 column (Pharmacia Uppsala, Sweden). Purification may be evaluated, e.g., by silver stained SDS-PAGE.

15 V. Preparation of antibodies against IL-171

Inbred Balb/c mice are immunized intraperitoneally, e.g., with 1 ml of purified human IL-171-FLAG emulsified in Freund's complete adjuvant on day 0, and in Freund's incomplete adjuvant on days 15 and 22. The mice are boosted with 0.5 ml of purified human IL-171 administered intravenously.

Polyclonal antiserum is collected. The serum can be purified to antibodies. The antibodies can be further processed, e.g., to Fab, Fab2, Fv, or similar fragments.

Hybridomas are created using, e.g., the non-secreting myeloma cells line SP2/0-Ag8 and polyethylene glycol 1000 (Sigma, St. Louis, MO) as the fusing agent. Hybridoma cells are placed in a 96-well Falcon tissue culture plate (Becton Dickinson, NJ) and fed with DMEM F12 (Gibco, Gaithersburg, MD) supplemented with 80 μg/ml gentamycin, 2 mM glutamine, 10% horse serum (Gibco, Gaithersburg, MD), 1% ADCM (CRTS, Lyon, France) 10<sup>-5</sup> M azaserine (Sigma, St. Louis, MO) and 5 x 10<sup>-5</sup> M hypoxanthine. Hybridoma supernatants are screened for antibody production against human IL-171 by immunocytochemistry (ICC) using acetone fixed human IL-171 transfected COS-7 cells and by ELISA using human IL-171-FLAG

35 transfected COS-7 cells and by ELISA using human IL-171-FLAG purified from COS-7 supernatants as a coating antigen.
Aliquots of positive cell clones are expanded for 6 days and

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cryopreserved as well as propagated in ascites from pristane (2,6,10,14-teramethylpentadecane, Sigma, St. Louis, MO) treated Balb/c mice who had received on intraperitoneal injection of pristane 15 days before. Typically, about 10<sup>5</sup> hybridoma cells in 1 ml of PBS are given intraperitoneally, and 10 days later, ascites are collected from each mouse.

After centrifugation of the ascites, the antibody fraction is isolated by ammonium sulfate precipitation and anion-exchange chromatography on a Zephyr-D silicium column (IBF Sepracor) equilibrated with 20 mM Tris pH 8.0. Proteins are eluted with a NaCl gradient (ranging from 0 to 1 M NaCl). 2 ml fractions are collected and tested by ELISA for the presence of anti-IL-171 antibody. The fractions containing specific anti-IL-171 activity are pooled, dialyzed, and frozen. Aliquots of the purified monoclonal antibodies may be peroxidase labeled.

Antibody preparations, polyclonal or monoclonal, may be cross absorbed, depleted, or combined to create reagents which exhibit desired combinations of selectivities and specificities. Defined specific antigens can be immobilized to a solid matrix and used to selectively deplete or select for desired binding capacities.

Similar methods will be applicable to IL-175, IL-172, IL-173, IL-174, IL-176, and/or IL-177. Methods to develop antibody preparations which cross react among various subsets of the family may be readily prepared.

#### VI. Quantification of human IL-171

Among the antibodies specific for IL-171, appropriate

clonal isolates are selected to quantitate levels of human IL
171 using a sandwich assay. Purified antibodies are diluted,
e.g., at 2 μg/ml in coating buffer (carbonate buffer, pH 9.6.

15 mM Na<sub>2</sub>CO<sub>3</sub>, 35 mM NaHCO<sub>3</sub>). This diluted solution is coated
onto the wells of a 96-well ELISA plate (Immunoplate Maxisorp

F96 certified, NUNC, Denmark) overnight at room temperature.
The plates are then washed manually, e.g., with a washing
buffer consisting of Phosphate Buffered Saline and 0.05% Tween

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20 (Technicon Diagnositics, USA). 110  $\mu$ l of purified human CTLA-8 diluted in TBS-B-T buffer [20 mM Tris, 150 mM NaCl, 18 BSA (Sigma, St. Louis, MO), and 0.05% Tween 20] is added to each well. After 3 hours of incubation at 37° C, the plates are washed once. 100  $\mu$ l of peroxidase labeled Ab16 diluted to 5  $\mu$ g/ml in TBS-B-T buffer is added to each well, and incubated for 2 hours at 37° C. The wells are then washed three times in washing buffer. 100  $\mu$ l of peroxidase substrate, 2.2' Azino-bis(3 ethylbenzthiazoine-6-sulfonic acid) (ABTS), diluted to 1 mg/ml in citrate/phosphate buffer, is added to each well, and the colorimetric reaction read at 405 nm.

### VII. Distribution of IL-170 genes.

The human IL-171 was identified from a sequence derived from an apoptotic T cell. The gene appears to be quite rare, which suggests the expression distribution would be highly restricted.

The human IL-175 was identified from a sequence derived from a 12 h thiouridine activated T cell. The gene appears to be quite rare, which suggests the expression distribution would be highly restricted.

The human IL-172 was identified from sequences derived from human fetal heart, liver and spleen, thymus, thymus tumor, and total fetus. Mouse was derived from sequences derived from mouse, embryo, mammary gland, and pooled organs. Both genes appear to be quite rare, which suggests their expression distribution would be highly restricted.

The human IL-173 was identified from sequence derived from a cDNA library from an epileptic brain frontal cortex. The rat IL-173 was derived from a cDNA library from cochlea, brain, cerebellum, eye, lung, and kidney. Again, the genes appear to be quite rare, which suggests the expression distributions would be highly restricted.

The mouse IL-174 was identified from sequence derived 35 from a cDNA library derived form a mouse embryo. The gene appears to be quite rare, which suggests the expression distribution would be highly restricted.

## VIII. Chromosome mapping of IL-170 genes

An isolated cDNA encoding the appropriate IL-170 gene is used. Chromosome mapping is a standard technique. See, e.g., BIOS Laboratories (New Haven, CT) and methods for using a mouse somatic cell hybrid panel with PCR.

The human IL-173 gene maps to human chromosome 13q11.

#### IX. Isolating IL-170 Homologues

A binding composition, e.g., antibody, is used for screening of an expression library made from a cell line which expresses an IL-170 protein. Standard staining techniques are used to detect or sort intracellular or surface expressed antigen, or surface expressing transformed cells are screened by panning. Screening of intracellular expression is performed by various staining or immunofluorescence procedures. See also McMahan, et al. (1991) EMBO J. 10:2821-2832.

Similar methods are applicable to isolate either

20 species or allelic variants. Species variants are isolated using cross-species hybridization techniques based upon a full length isolate or fragment from one species as a probe, or appropriate species.

#### 25 X. Isolating receptors for IL-170

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Methods are available for screening of an expression library made from a cell line which expresses potential IL-170 receptors. A labeled IL-170 ligand is produced, as described above. Standard staining techniques are used to detect or sort surface expressed receptor, or surface expressing transformed cells are screened by panning. See also McMahan, et al. (1991) <u>EMBO J.</u> 10:2821-2832.

For example, on day 0, precoat 2-chamber permanox slides with 1 ml per chamber of fibronectin, 10 ng/ml in PBS, for 30 min at room temperature. Rinse once with PBS. Then plate COS cells at  $2-3 \times 10^5$  cells per chamber in 1.5 ml of growth media. Incubate overnight at  $37^{\circ}$  C.

On day 1 for each sample, prepare 0.5 ml of a solution of 66 µg/ml DEAE-dextran, 66 µM chloroquine, and 4 µg DNA in serum free DME. For each set, a positive control is prepared, e.g., of huIL-170-FLAG cDNA at 1 and 1/200 dilution, and a negative mock. Rinse cells with serum free DME. Add the DNA solution and incubate 5 hr at 37° C. Remove the medium and add 0.5 ml 10% DMSO in DME for 2.5 min. Remove and wash once with DME. Add 1.5 ml growth medium and incubate overnight.

10 On day 2, change the medium. On days 3 or 4, the cells are fixed and stained. Rinse the cells twice with Hank's Buffered Saline Solution (HBSS) and fix in 4% paraformaldehyde (PFA)/glucose for 5 min. Wash 3X with The slides may be stored at -80° C after all liquid is removed. For each chamber, 0.5 ml incubations are performed as follows. Add HBSS/saponin (0.1%) with 32 µ1/ml of 1 M NaN3 for 20 min. Cells are then washed with HBSS/saponin 1X. Soluble antibody is added to cells and incubate for 30 min. Wash cells twice with HBSS/saponin. Add second antibody, e.g., Vector anti-mouse antibody, at 20 1/200 dilution, and incubate for 30 min. Prepare ELISA solution, e.g., Vector Elite ABC horseradish peroxidase solution, and preincubate for 30 min. Use, e.g., 1 drop of solution A (avidin) and 1 drop solution B (biotin) per 2.5 25 ml HBSS/saponin. Wash cells twice with HBSS/saponin. ABC HRP solution and incubate for 30 min. Wash cells twice with HBSS, second wash for 2 min, which closes cells. add Vector diaminobenzoic acid (DAB) for 5 to 10 min. drops of buffer plus 4 drops DAB plus 2 drops of H2O2 per 5 30 ml of glass distilled water. Carefully remove chamber and rinse slide in water. Air dry for a few minutes, then add 1 drop of Crystal Mount and a cover slip. Bake for 5 min at 85-90° C.

Alternatively, the labeled ligand is used to affinity
35 purify or sort out cells expressing the receptor. See,
e.g., Sambrook, et al. or Ausubel, et al.

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All references cited herein are incorporated herein by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

Many modifications and variations of this invention can be made without departing from its spirit and scope, as will be apparent to those skilled in the art. The specific embodiments described herein are offered by way of example only, and the invention is to be limited only by the terms of the appended claims, along with the full scope of equivalents to which such claims are entitled.

# WHAT IS CLAIMED IS:

- 1. An isolated or recombinant polynucleotide comprising sequence from a mammalian IL-171, which:
- 5 a) encodes at least 8 contiguous amino acids of SEQ ID NO: 3 or 5;
  - b) encodes at least two distinct segments of at least 5 contiguous amino acids of SEQ ID NO: 3 or 5; or
  - c) comprises one or more segments of at least 21 contiguous nucleotides of SEQ ID NO: 1 or 4.
  - 2. The polynucleotide of Claim 1 in an expression vector, comprising a sequence which:
  - a) encodes at least 12 contiguous amino acids of SEQ ID
     NO: 3 or 5;
    - b) encodes at least two distinct segments of at least 7 and 10 contiguous amino acids of SEQ ID NO: 3 or 5; or
    - c) comprises at least 27 contiguous nucleotides of SEQ ID NO: 1 or 4.
  - 3. The polynucleotide of Claim 2 which:
    - a) encodes at least 16 contiguous amino acids of SEQ IDNO: 3 or 5;
- 25 b) encodes at least two distinct segments of at least 10 and 14 contiguous amino acid residues of SEQ ID NO: 3 or 5; or
  - c) comprises at least 33 contiguous nucleotides of SEQ ID NO: 1 or 4.

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- 4. A method of making:
  - a) a polypeptide comprising expressing said expression vector of Claim 2, thereby producing said polypeptide;
- 35 b) a duplex nucleic acid comprising contacting a polynucleotide of Claim 2 with a complementary

nucleic acid, thereby resulting in production of said duplex nucleic acid; or

c) a polynucleotide of Claim 2 comprising amplifying using a PCR method.

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- 5. An isolated or recombinant polynucleotide which hybridizes under stringent wash conditions of at least 55° C and less than 400 mM salt to:
- a) the (IL-171) polynucleotide of Claim 3 which consists

  of the entire mature coding portions of SEQ ID NO: 1

  or 4.
  - 6. A polynucleotide of Claim 5:
    - a) wherein said wash conditions are at least 65° C and less than 300 mM salt; or
    - b) which comprises at least 50 contiguous nucleotides of the coding portion of SEQ ID NO: 1 or 4.
  - 7. A kit comprising said polynucleotide of Claim 6, and
- 20 a) instructions for the use of said polynucleotide for detection;
  - b) instructions for the disposal of said polynucleotide or other reagents of said kit; or
  - c) both a and b.

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- 8. A cell containing said expression vector of Claim 3, wherein said cell is:
  - a) a prokaryotic cell;
  - b) a eukaryotic cell;
- 30 c) a bacterial cell;
  - d) a yeast cell;
  - e) an insect cell;
  - f) a mammalian cell;
  - g) a mouse cell;
- 35 h) a primate cell; or
  - i) a human cell.

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- 9. An isolated or recombinant antigenic polypeptide comprising at least:
  - a) one segment of 8 identical contiguous amino acids from SEQ ID NO: 3 or 5; or
- 5 b) two distinct segments of at least 5 contiguous amino acids from SEQ ID NO: 3 or 5.
  - 10. The polypeptide of Claim 9, wherein:
    - a) said segment of 8 identical contiguous amino acids is at least 14 contiguous amino acids; or
    - b) one of said segments of at least 5 contiguous amino acids comprises at least 7 contiguous amino acids.
- 11. The polypeptide of Claim 9, wherein said 15 polypeptide:
  - a) comprises a mature sequence of SEQ ID NO: 3 or 5;
  - b) binds with selectivity to a polyclonal antibody generated against an immunogen of SEQ ID NO: 3 or 5;
  - c) comprises a plurality of distinct polypeptide segments of 10 contiguous amino acids of SEQ ID NO: 3 or 5;
    - d) is a natural allelic variant of SEQ ID NO: 3 or 5;
    - e) has a length at least 30 amino acids; or
    - f) exhibits at least two non-overlapping epitopes which are selective for SEQ ID NO: 3 or 5.
    - 12. The polypeptide of Claim 11, which:
      - a) is in a sterile composition;
      - b) is not glycosylated;
- 30 c) is denatured;
  - d) is a synthetic polypeptide;
  - e) is attached to a solid substrate;
  - f) is a fusion protein with a detection or purification tag;
- 35 g) is a 5-fold or less substitution from a natural sequence; or

- h) is a deletion or insertion variant from a natural sequence.
- 13. A method using said polypeptide of Claim 9:
- 5 a) to label said polypeptide, comprising labeling said polypeptide with a radioactive label;
  - b) to separate said polypeptide from another polypeptide in a mixture, comprising running said mixture on a chromatography matrix, thereby separating said polypeptides;
  - c) to identify a compound that binds selectively to said polypeptide, comprising incubating said compound with said polypeptide under appropriate conditions; thereby causing said compound to bind to said polypeptide; or
  - d) to conjugate said polypeptide to a matrix, comprising derivatizing said polypeptide with a reactive reagent, and conjugating said polypeptide to said matrix.

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14. A binding compound comprising an antigen binding portion from an antibody which binds with selectivity to said polypeptide of Claim 11, wherein said polypeptide comprises SEQ ID NO 3 or 5.

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- 15. The binding compound of Claim 14, wherein said antibody is a polyclonal antibody which is raised against SEQ ID NO: 3 or 5.
- 30 16. The binding compound of Claim 14, wherein said:
  - a) antibody:
    - i) is immunoselected;
    - ii) binds to a denatured protein; or
    - iii) exhibits a Kd to said polypeptide of at least
      30 mM; or
  - b) said binding compound:

- i) is attached to a solid substrate, including a bead or plastic membrane;
- ii) is in a sterile composition; or
- iii) is detectably labeled, including a radioactive or fluorescent label.
- 17. A method of producing an antigen:antibody complex, comprising contacting a polypeptide comprising sequence from SEQ ID NO: 3 or 5 with a binding compound of Claim 14 under conditions which allow said complex to form.
- 18. The method of Claim 17, wherein said binding compound is an antibody, and said polypeptide is in a biological sample.

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- 19. A kit comprising said binding compound of Claim 14 and:
  - a) a polypeptide of SEQ ID NO: 3 or 5;
  - b) instructions for the use of said binding compound for detection; or
  - c) instructions for the disposal of said binding compound or other reagents of said kit.
- 20. A method of evaluating the selectivity of binding of an antibody to a protein of SEQ ID NO: 3 or 5, comprising contacting said antibody to said protein and to another cytokine; and comparing binding of said antibody to said protein and said cytokine.

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